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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT) (51) International Patent Classification 6: (11) International Publication Number: WO 97/26277 C07K 14/51, G06F 17/50 A2 (43) International Publication Date: 24 July 1997 (24.07.97) (21) International Application Number: PCT/US97/01071 (81) Designated States: AU, CA, JP, European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, (22) International Filing Date: 22 January 1997 (22.01.97) PT, SE). (30) Priority Data: Published 08/589,552 Without international search report and to be republished 22 January 1996 (22.01.96) US upon receipt of that report. (71) Applicants: CREATIVE BIOMOLECULES, INC. [US/US]; 45 South Street, Hopkinton, MA 01748 (US). BRANDEIS UNIVERSITY [US/US]; 45 South Street, Waltham, MA 02254-9110 (US). (72) Inventors: KECK, Peter; 50 Dolan Road, Millbury, MA 01527 (US). GRIFFITH, Diana, L.; 15 Woodridge Circle, Weston, MA 02193 (US). CARLSON, William, D.; 40 Black Oak Road, Weston, MA 02193 (US). RUEGER, David, C.; 19 Downey Street, Hopkinton, MA 01748 (US). SAMPATH, Kuber, T.; 6 Spring Street, Medway, MA 02053 (US). (74) Agent: GREENHALGH, Duncan, A.; Testa, Hurwitz & Thibeault, L.L.P., High Street Tower, 125 High Street, Boston, MA 02110 (US).

#### (54) Title: METHODS AND COMPOSITIONS FOR PRODUCING MORPHOGEN ANALOGS

#### (57) Abstract

The invention disclosed herein provides methods and compositions for the computer-assisted design of morphogen analogs. Practice of the invention is enabled by the use of at least a portion of the atomic co-ordinates defining the three-dimensional structure of human osteogenic protein-1 (hOP-1) as a starting point in the design of the morphogen analogs. In addition, the invention provides methods for producing morphogen analogs of interest, and methods for testing whether the resulting analogs mimic or agonize human OP-1-like biological activity. The invention also provides a family of morphogen analogs produced by such methods.

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# METHODS AND COMPOSITIONS FOR PRODUCING MORPHOGEN ANALOGS

#### Related Applications

This application is a continuation-in-part of copending application U.S.S.N. 08/589,552, filed January 22, 1996, the disclosure of which is incorporated herein by reference.

#### Field of the Invention.

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The present invention relates generally to methods and compositions for designing, identifying, and producing compounds useful as tissue morphogenic protein analogs. More specifically, the invention relates to structure-based methods and compositions useful in designing, identifying, and producing molecules which act as functional mimetics of the tissue morphogenic protein osteogenic protein-1 (OP-1).

#### Background of the Invention

Cell differentiation is the central characteristic of tissue morphogenesis which initiates during embryogenesis, and continues to various degrees throughout the life of an organism in adult tissue repair and regeneration mechanisms. The degree of morphogenesis in adult tissue varies among different tissues and is related, among other things, to the degree of cell turnover in a given tissue.

The cellular and molecular events which govern the stimulus for differentiation of cells is an area of intensive research. In the medical and veterinary fields, it is anticipated that discovery of the factor or factors which control cell differentiation and tissue morphogenesis will advance significantly the ability to repair and regenerate diseased or damaged mammalian tissues and organs. Particularly useful areas for human and veterinary therapeutics include reconstructive surgery, the treatment of tissue degenerative diseases including, for example, arthritis, emphysema, osteoporosis, cardiomyopathy, cirrhosis, degenerative nerve diseases, inflammatory diseases, and cancer, and in the regeneration of tissues, organs and limbs. In this and related applications, the terms "morphogenetic" and "morphogenic" are used interchangeably.

A number of different factors have been isolated in recent years which appear to play a role in cell differentiation. Recently, a distinct subfamily of the "superfamily" of structurally related proteins referred to in the art as the "transforming growth factor-  $\beta$  (TGF- $\beta$ ) superfamily of proteins" have been identified as true tissue morphogens.

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The members of this distinct "subfamily" of true tissue morphogenic proteins share substantial amino acid sequence homology within their morphogenetically active C-terminal domains (at least 50% identity in the C-terminal 102 amino acid sequence), including a conserved six or seven cysteine skeleton, and share the *in vivo* activity of inducing tissue-specific morphogenesis in a variety of organs and tissues. The proteins apparently contact and interact with progenitor cells e.g., by binding suitable cell surface molecules, predisposing or otherwise stimulating the cells to proliferate and differentiate in a morphogenetically permissive environment. These morphogenic proteins are capable of inducing the developmental cascade of cellular and molecular events that culminate in the formation of new organ-specific tissue, including any vascularization, connective tissue formation, and nerve innervation as required by the naturally occurring tissue. The proteins have been shown to induce morphogenesis of both bone cartilage and bone, as well as periodontal tissues, dentin, liver, and neural tissue, including retinal tissue.

True tissue morphogenic proteins identified to date include proteins originally identified as bone inductive proteins. These include OP-1, (osteogenic protein-1, also referred to in related applications as "OP1"), its Drosophila homolog, 60A, with which it shares 69% identity in the C-terminal "seven cysteine" domain, and the related proteins OP-2 (also referred to in related applications as "OP2") and OP-3, both of which share approximately 65-75% identity with OP-1 in the C-terminal seven cysteine domain, as well as BMP5, BMP6 and its murine homolog, Vgr-1, all of which share greater than 85% identity with OP-1 in the C-terminal seven cysteine domain, and the BMP6 Xenopus homolog, Vgl, which shares approximately 57% identity with OP-1 in the C-terminal seven cysteine domain. Other bone inductive proteins include the CBMP2 proteins (also referred to in the art as BMP2 and BMP4) and their Drosophila homolog, DPP. Another tissue morphogenic protein is GDF-1 (from mouse). See, for example, PCT documents US92/01968 and US92/07358, the disclosures of which are incorporated herein by reference. Members of the BMP/OP subfamily and the amino acid sequence identities

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(expressed as percentages) between selected members of the TGF-B superfamily are shown in Fig. 6.

As stated above, these true tissue morphogenic proteins are recognized in the art as a distinct subfamily of proteins different from other members of the TGF-B superfamily in that they share a high degree of sequence identity in the C-terminal domain and in that the true tissue morphogenic proteins are able to induce, on their own, the full cascade of events that result in formation of functional tissue rather than merely inducing formation of fibrotic (scar) tissue. Specifically, members of the family of morphogenic proteins are capable of all of the following in a morphogenetically permissive environment: stimulating cell proliferation and cell differentiation, and supporting the growth and maintenance of differentiated cells. The morphogenic proteins apparently also may act as endocrine, paracrine or autocrine factors.

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The morphogenic proteins are capable of significant species "crosstalk." That is, xenogenic (foreign species) homologs of these proteins can substitute for one another in functional activity. For example, dpp and 60A, two Drosophila proteins, can substitute for their mammalian homologs, BMP2/4 and OP-1, respectively, and induce endochondral bone formation at a non-bony site in a standard rat bone formation assay. Similarly, BMP2 has been shown to rescue a dpp<sup>-</sup> mutation in Drosophila. In their native form, however, the proteins appear to be tissue-specific, each protein typically being expressed in or provided to one or only a few tissues or, alternatively, expressed only at particular times during development. For example, GDF-1 appears to be expressed primarily in neural tissue, while OP-2 appears to be expressed at relatively high levels in early (e.g., 8-day) mouse embryos. The endogenous morphogens may be synthesized by the cells on which they act, by neighboring cells, or by cells of a distant tissue, the secreted protein being transported to the cells to be acted on.

A particularly potent tissue morphogenic protein is OP-1. This protein, and its xenogenic homologs, are expressed in a number of tissues, primarily in tissues of urogenital origin, as well as in bone, mammary and salivary gland tissue, reproductive tissues, and gastrointestinal tract tissue. It is expressed also in different tissues during embryogenesis, its presence coincident with the onset of morphogenesis of that tissue.

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The morphogenic protein signal transduction across a cell membrane appears to occur as a result of specific binding interaction with one or more cell surface receptors. Recent studies on cell surface receptor binding of various members of the TGF-ß protein superfamily suggests that the ligands mediate their activity by interaction with two different receptors, referred to as Type I and Type II receptors to form a hetero-complex. A cell surface bound beta-glycan also may enhance the binding interaction. The Type I and Type II receptors are both serine /threonine kinases, and share similar structures: an intracellular domain that consists essentially of the kinase, a short, extended hydrophobic sequence sufficient to span the membrane one time, and an extracellular domain characterized by a high concentration of conserved cysteines.

Morphogenic proteins are disulfide-linked dimers which are expressed as large precursor polypeptide chains containing a hydrophobic signal sequence, a long and relatively poorly conserved N-terminal pro region of several hundred amino acids, a cleavage site and a mature domain comprising an N-terminal region which varies among the family members and a more highly conserved C-terminal region. The C-terminal region, which is present in the processed mature proteins of all known morphogen family members, contains approximately 100 amino acids with a characteristic motif having a conserved six or seven cysteine skeleton. Each of the morphogenic proteins isolated to date are dimeric structures wherein the monomer subunits are held together by non-covalent interactions or by one or more disulfide bonds. The morphogenic proteins are active as dimeric proteins but are inactive as individual monomer subunits.

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As a result of their biological activities, significant effort has been directed toward the development of morphogen-based therapeutics for treating injured or diseased mammalian tissue, including, for example, therapeutic compositions for inducing regenerative healing of bone defects such as fractures, as well as therapeutic compositions for preserving or restoring healthy metabolic properties in diseased bone tissue, e.g., osteopenic bone tissue. Complete descriptions of efforts to develop and characterize morphogen-based therapeutics for non-chondrogenic tissue applications in mammals, particularly humans, are set forth, for example, in: EP 0575,555; WO93/04692; WO93/05751; WO94/06399; WO94/03200; WO94/06449; WO94/10203; and WO94/06420, the disclosures of each of which are incorporated herein by reference.

Certain difficulties may be experienced upon administration of naturally isolated or recombinantly produced morphogenic proteins to a mammal. These difficulties may include, for

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example, loss of morphogenic activity due to disassociation of the biologically active morphogen dimer into its inactive monomer subunits, and/or handling problems due to low solubility under physiological conditions.

Accordingly, a need remains for the identification of morphogen analogs, which mimic or enhance the physiological effects of a morphogenic protein, for example OP-1. The analogs may be modified, morphogenically active hOP-1 protein dimers, or fragments or truncated analogs thereof, peptides or small organic molecules. Preferably the analogs have enhanced therapeutic value, for example, by being more stable and/or more soluble under physiological conditions than naturally occurring hOP-1, or, for example, by having enhanced tissue targeting specificity, enhanced biodistribution or a reduced clearance rate in the body.

It is an object of the present invention to provide a database defining the atomic coordinates of the three-dimensional structure of mature hOP-1, all or a portion of which can be
used as part of a computer system for designing and/or identifying a functional analog of hOP-1.

Another object is to provide means for designing and/or identifying a molecule having enhanced
solubility and/or stability under physiological conditions as compared with hOP-1 and which is
capable of mimicking or enhancing the biological activity of hOP-1 in a mammal. Another
object of the invention is to provide a therapeutic composition comprising an analog designed
and/or identified, and produced by the methods of the invention, and suitable for administration
to a mammal in need thereof, such as a mammal afflicted with a metabolic bone disease, e.g., a
disease characterized by osteopenia. Another object of the invention is to provide methods and
compositions useful for designing and/or identifying, and producing an hOP-1 antagonist capable
of, for example, competing with hOP-1 for receptor binding, but incapable of inducing a
receptor-mediated downstream biological effect.

These and other objects and features of the invention will be apparent from the description, drawings, and claims which follow.

#### Summary of the Invention

The present invention is based, in part, upon the X-ray crystallographic determination of the three-dimensional structure of mature, dimeric human osteogenic protein-1 (hOP-1). The three-dimensional structure of hOP-1 has been resolved to 2.3Å. Provided herein are two sets of

atomic X-ray crystallographic co-ordinates for hOP-1, one set defining a hOP-1 structure resolved to a resolution of 2.8Å, and the other set defining a hOP-1 structure resolved to a resolution of 2.3Å. With this disclosure, the skilled artisan is provided with sets of atomic co-ordinates for use in conventional computer aided design (CAD) methodologies to identify or design protein or peptide analogs of OP-1, or alternatively, to identify or design small organic molecules that functionally mimic OP-1.

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In one aspect, the invention provides a computer system comprising a memory and a processor in electrical communication with the memory. The memory has disposed therein, atomic X-ray crystallographic co-ordinates which together define at least a portion of the three-dimensional structure of hOP-1. In a preferred embodiment, the atomic co-ordinates are defined by either a portion or all of the atomic co-ordinates set forth in Figure 15 or Figure 16.

The processor, in electrical communication with the memory, comprises a process which generates a molecular model having a three-dimensional shape representative of at least a portion of human OP-1. In a preferred embodiment, the processor is capable of producing a molecular model having, in addition to the three-dimensional shape, a solvent accessible surface representative of at least a portion of human OP-1.

As used herein, the term "computer system" is understood to mean any general or special purpose system which includes a processor in electrical communication with both a memory and at least one input/output device, such as a terminal. Such a system may include, but is not limited to, personal computers, workstations or mainframes. The processor may be a general purpose processor or microprocessor or a specialized processor executing programs located in RAM memory. The programs may be placed in RAM from a storage device, such as a disk or preprogrammed ROM memory. The RAM memory in one embodiment is used both for data storage and program execution. The term computer system also embraces systems where the processor and memory reside in different physical entities but which are in electrical communication by means of a network.

In the present invention, the processor executes a modeling program which accesses data representative of the X-ray crystallographic co-ordinates of hOP-1 thereby to construct a three-dimensional model of the molecule. In addition, the processor also can execute another program,

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a solvent accessible surface program, which uses the three-dimensional model of hOP-1 to construct a solvent accessible surface of at least a portion of the hOP-1 molecule and optionally calculate the solvent accessible areas of atoms. In one embodiment the solvent accessible surface program and the modeling program are the same program. In another embodiment, the modeling program and the solvent accessible surface program are different programs. In such an embodiment the modeling program may either store the three-dimensional model of hOP-1 in a region of memory accessible both to it and to the solvent accessible surface program, or the three-dimensional model may be written to external storage, such as a disk, CD ROM, or magnetic tape for later access by the solvent accessible surface program.

The memory may have stored therein the entire set of X-ray crystallographic co-ordinates which define mature biologically active human OP-1, or may comprise a subset of such co-ordinates including, for example, one or more of: a finger 1 region; a finger 2 region; and a heel region. The protein structures which correspond to the finger and heel regions are described in detail below.

In another preferred embodiment, the processor also is capable of identifying a morphogen analog, or a morphogen antagonist for example, a protein, peptide or small organic molecule, having a three-dimensional shape and preferably, in addition, a solvent accessible surface corresponding to at least a portion of human OP-1 and competent to mimic an OP-1 specific activity.

As used herein, with respect to OP-1 (or related morphogens), or with respect to a region of OP-1, the phrase "at least a portion of the three-dimensional structure of" or "at least a portion of" is understood to mean a portion of the three-dimensional surface structure of the morphogen, or region of the morphogen, including charge distribution and hydrophilicity/hydrophobicity characteristics, formed by at least three, more preferably at least three to ten, and most preferably at least ten contiguous amino acid residues of the OP-1 monomer or dimer. The contiguous residues forming such a portion may be residues which form a contiguous portion of the primary structure of the OP-1 molecule, residues which form a contiguous portion of the three-dimensional surface of the OP-1 monomer, residues which form a contiguous portion of the three-dimensional surface of the OP-1 dimer, or a combination thereof. Thus, the residues forming a portion of the three-dimensional structure of OP-1 need not be contiguous in the

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primary sequence of the morphogen but, rather, must form a contiguous portion of the surface of the morphogen monomer or dimer. In particular, such residues may be non-contiguous in the primary structure of a single morphogen monomer or may comprise residues from different monomers in the dimeric form of the morphogen. As used herein, the residues forming "a portion of the three-dimensional structure of " a morphogen, or "a portion of" a morphogen, form a contiguous three-dimensional surface in which each atom or functional group forming the portion of the surface is separated from the nearest atom or functional group forming the portion of the surface by no more than 40 Å, preferably by no more than 20 Å, more preferably by no more than 5-10 Å, and most preferably by no more than 1-5 Å.

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As used herein the term "X-ray crystallographic co-ordinates" refers to a series of mathematical co-ordinates (represented as "X", "Y" and "Z" values) that relate to the spatial distribution of reflections produced by the diffraction of a monochromatic beam of X-rays by atoms of an hOP-1 molecule in crystal form. The diffraction data are used to generate electron density maps of the repeating units of a crystal, and the resulting electron density maps are used to define the positions of individual atoms within the unit cell of the crystal.

As will be apparent to those of ordinary skill in the art, the hOP-1 structure presented herein is independent of its orientation, and that the atomic co-ordinates listed in Figs. 15 and 16 merely represent one possible orientation of the hOP-1 structure. It is apparent, therefore, that the atomic co-ordinates listed in Figs. 15 and 16, may be mathematically rotated, translated, scaled, or a combination thereof, without changing the relative positions of atoms or features of the hOP-1 structure. Such mathematical manipulations are intended to be embraced herein. Furthermore, it will be apparent to the skilled artisan that the X-ray atomic co-ordinates defined herein have some degree of uncertainty in location (see, for example, column "δ" in Fig. 16 which shows the thermal uncertainty in location of each atom, as expressed in Å). Accordingly, for purposes of this invention, a preselected protein or peptide having the same amino acid sequence as at least a portion of hOP-1 is considered to have the same structure as the corresponding portion of hOP-1, when a set of atomic co-ordinates defining backbone Cα atoms of the preselected protein or peptide can be superimposed onto the corresponding Cα atoms for hOP-1 (as listed in Figure 16) to a root mean square deviation of preferably less than about 1.5 Å, and most preferably less than about 0.75 Å.

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As used herein, the term "morphogen analog", is understood to mean any molecule capable of mimicking OP-1's receptor binding activity and/or and inducing a receptor mediated downstream biological effect characteristic of a morphogenic protein. Inducing alkaline phosphatase activity is a characteristic biological effect. The analog may be a protein, peptide, or non-peptidyl based organic molecule. Accordingly, the term morphogen analog embraces any substance having such OP-1 like activity, regardless of the chemical or biochemical nature thereof. The present morphogen analog can be a simple or complex substance produced by a living system or through chemical or biochemical synthetic techniques. It can be a large molecule, e.g., a modified hOP-1 dimer produced by recombinant DNA methodologies, or a small molecule, e.g., an organic molecule prepared *de novo* according to the principles of rational drug design. It can be a substance which is a mutein (or mutant protein) of hOP-1, a substance that structurally resembles a solvent-exposed surface epitope of hOP-1 and binds an OP-1 specific receptor, or a substance that otherwise stimulates an OP-1 specific receptor displayed on the surface of an OP-1 responsive cell.

As used herein, the terms "OP-1 or OP-1-like biological activity" are understood to mean any biological activities known to be induced or enhanced by OP-1. OP-1 and OP-1-like biological activities include, but are not limited to, stimulating proliferation of progenitor cells; stimulating differentiation of progenitor cells; stimulating proliferation of differentiated cells; and supporting growth and maintenance of differentiated cells. The term "progenitor cells" includes uncommitted cells, preferably of mammalian origin that are competent to differentiate into one or more specific types of differentiated cells, depending on their genomic repertoire and the tissue specificity of the permissive environment where morphogenesis is induced. Specifically, with regard to bone, cartilage, nerve, and liver tissue, the OP-1 stimulated morphogenic cascade culminates in the formation of new or regenerative differentiated tissue appropriate to the selected local environment. OP-1 mediated morphogenesis, therefore, differs significantly from simple reparative healing processes in which scar tissue (e.g., fibrous connective tissue) is formed and fills a lesion or other defect in differentiated functional tissue.

As used herein a "morphogen antagonist" is a molecule competent to mimic OP-1 receptor binding activity but which cannot induce a receptor-mediated downstream effect.

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In yet another preferred embodiment, the processor is capable of identifying amino acids defined by the co-ordinates, which upon site-directed modification, either by chemical modification or amino acid substitution, enhance the solubility and/or stability of human OP-1.

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In a related aspect, the invention provides a method of producing a morphogen analog that mimics or enhances an OP-1 or OP-1-like biological activity. The method comprises the steps of: (a) providing a molecular model defining a three-dimensional shape representative of at least a portion of human OP-1, (b) identifying a compound having a three-dimensional shape corresponding to the three-dimensional shape representative of at least the portion of human OP-1; and (c) producing the compound identified in step (b). The method can comprise the additional step of testing the compound in a biological system to determine whether the resultant candidate compound mimics or agonizes the biological activity of OP-1. It is contemplated that, in the aforementioned method, step (a) and/or (b) may be performed by means of an electronic processor using commercially available software packages.

It is contemplated that, upon determination of whether the candidate compound modulates OP-1 activity, the candidate compound can be iteratively improved using conventional CAD and/or rational drug design methodologies, well known and thoroughly documented in the art. Furthermore, it is contemplated that the resultant compound identified thus far, may be produced in a commercially useful quantity for administration into a mammal.

In another embodiment, the morphogen analog is created using atomic co-ordinates set forth in either Figs. 15 or 16. By reviewing the atomic co-ordinates set forth in Figs. 15 and 16, the skilled artisan can observe the three-dimensional structure of particular amino acid sequences located *in situ* within the three-dimensional structure of hOP-1. Preferred amino acid sequences are defined by one or more of the peptides selected from the group consisting of: H1, H-n2, H-c2, F1-2, F2-2 and F2-3, as discussed hereinbelow. The peptides provide templates which can be used in the production of more effective morphogen analogs. In a preferred embodiment, the  $C\alpha$  atoms of amino acid residues in the morphogen analog are located within 6Å, preferably within 3Å, and most preferably within 2Å of the corresponding  $C\alpha$  atom as defined by the respective atomic co-ordinates in Figs. 15 or 16. In another preferred embodiment, the  $C\alpha$  atoms of amino acid residues in the morphogen analog are located within 6Å, preferably within 3Å, and most preferably within 2Å of the corresponding  $C\alpha$  atoms of at least three amino acids in the peptide

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sequences H1, H-n2, H-c2, F1-2, F2-2 and F2-3, wherein each of the Cα atoms in the peptides are defined by the respective atomic co-ordinates set forth in Figs. 15 or 16.

In another embodiment, the invention provides morphogen analogs having greater solubility and/or stability in aqueous buffers than native dimeric hOP-1. In yet another embodiment, the invention provides a morphogen analog which is a modified form of dimeric hOP-1, in which the modification eliminates an epitope or region on OP-1 normally recognized by an antibody or by a cellular scavenging protein for clearing OP-1 from the body.

In another embodiment, the invention provides means for creating an analog with altered receptor binding characteristics. For example, provided with the structure, charge distribution, and solvent accessible surface information pertaining to the putative receptor binding site, one can alter or modify receptor binding specificity and avidity. In one embodiment, amino acid replacements in this region are made with reference to the corresponding amino acids of other known morphogens, disclosed for example, in WO 94/06449 or WO 93/05751.

After having determined the three-dimensional structure of human OP-1, a skilled artisan, in possession of the atomic co-ordinates defining the OP-1 structure is hereby enabled to use conventional CAD and/or rational drug design methodologies to identify or design protein or peptide analogs, or other small organic molecules which, after having been produced using conventional chemistries and methodologies, can be tested either *in vitro* or *in vivo* to assess whether they mimic or enhance the biological activity of human OP-1.

The foregoing and other objects, features and advantages of the present invention will be made more apparent from the following detailed description of preferred embodiments of the invention.

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#### Brief Description of the Drawings.

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The file of this patent contains at least one drawing executed in color. Copies of this patent with color drawing(s) will be provided by the Patent and Trademark Office upon request and payment of the necessary fee.

The objects and features of the invention may be better understood by reference to the drawings described below, wherein like referenced features identify common features in corresponding figures.

Figure 1A is a simplified line drawing useful in describing the structure of a monomeric subunit of hOP-1. See the Summary of the Invention, *infra*, for explanation. Figures 1B, 1C, and 1D are monovision ribbon tracings of the respective peptide backbones of hOP-1 finger-1, heel, and finger-2 regions. Figures 1E and 1F are schematic representations of monomeric and dimeric forms of hOP-1, respectively, as represented by a left hand motif.

Fig. 2 is a schematic drawing of a monomeric subunit of hOP-1. The hOP-1 cysteine knot comprising three disulfide bonds constitutes the core of the hOP-1 monomer subunit. Two disulfide bonds which connect residues Cys 67 - Cys 136 and Cys 71-Cys 138 produce an eight residue ring through which the third disulfide bond which connects residues Cys 38 - Cys 104 passes. Four strands of antiparallel B-sheet, which emanate from the knot, form the two finger like projections. An a-helix located on the opposite end of the knot, lies perpendicular to the axis of the two fingers thereby forming the heel. The N-terminus of the monomer subunit remains unresolved. The \( \beta \)-sheets are displayed as arrows and labeled from \( \beta 1 \) through \( \beta 8 \). The α-helix is displayed as a tube and labeled ∝l. The intra-subunit disulfide bonds that constitute the cysteine knot are shown in solid lines. Starting from Gln 36 ("N<sub>36</sub>"), the first residue shown in this figure, the amino acid residues which produce secondary structure in the Finger 1 region include: Lys 39 - His 41 (B1), Tyr 44-Ser 46 (B2), Glu 60 - Ala 63 (B3), Tyr 65 - Glu 70 (B4); the amino acid residues which produce secondary structure in the Finger 2 region include: Cys 103-Asn 110 (B5); Ile 112 - Asp 118 (B6); Asn 122 - Tyr 128 (B7); Val 132 - His 139 (B8); and the amino acid residues which produce secondary structure in the heel region include: Thr 82 - Ile 94(a1).

Figure 3 is a structure-based sequence alignment of the hOP-1 and TGF-ß2 finger-1, heel, and finger-2 regions. Amino acid residues in the heel regions which constitute inter-chain contacts in the dimers of hOP-1 and TGF-ß2 are highlighted as white on black. Amino acid residues in the finger-1 and finger-2 regions which contact the other chain are highlighted as black on gray. In hOP-1 and TGF-ß2, the amino acids located at the same residue positions constitute the inter-chain contacts.

Figures 4A and 4B are stereo peptide backbone ribbon trace drawings illustrating the three-dimensional shape of hOP-1: A) from the "top" (down the two-fold axis of symmetry between the subunits) with the axes of the helical heel regions generally normal to the paper and the axes of each of the finger 1 and finger 2 regions generally vertical, and B) from the "side" with the two-fold axis between the subunits in the plane of the paper, with the axes of the heels generally horizontal, and the axes of the fingers generally vertical. The hOP-1 monomer has an accessible non-polar surface area of approximately 4394Ų, while that for the dimer is approximately 6831Ų resulting in a hidden area upon dimerization of approximately 979Ų per monomer. The reader is encouraged to view the stereo alpha carbon trace drawings in wall-eyed stereo, for example, using a standard stereo viewer device, to more readily visualize the spatial relationships of amino acids sequences in the morphogen analog design.

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Figure. 5A is a backbone ribbon trace drawing illustrating the hOP-1 dimer comprising the two hOP-1 monomer subunits resolved to 2.8Å. One monomer subunit is shown in green and the other monomer subunit is shown in gold. Amino acid residues disposed within the purported receptor binding domain having solvent accessible side chains are shown as atomic spheres. The tips of the finger 1 and finger 2 regions of one OP-1 monomeric subunit and a loop at the C-terminal end of the heel of the other OP-1 monomeric subunit are believed to constitute the receptor binding domain. Amino acids located at positions of variable amino acid sequence shown in white while amino acids located at more conserved positions are shown in red. Figures 5B and 5C are pictures showing the respective solvent accessible surfaces of OP-1 and TGF-B2 dimers colored based on their electrostatic potential. Surface regions having an electrostatic potential of -3 kT or less are shown in red while surface regions of +3 kT or greater are shown in blue. Neutral regions are shown in green or gold to correspond to the backbone ribbons shown in 5A.

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Figure 6 is a table showing an identity matrix for the TGF-B superfamily. The matrix comprises members of the TGF-B superfamily having an amino acid sequence identity relative to OP-1 of greater than 36%. In the matrix, the TGF- B superfamily members are placed in order of decreasing amino acid identity relative to OP-1. TGF- B2 has an amino acid sequence of identity of 36% relative to OP-1 and is positioned the bottom of the matrix. Boxes enclose families of sequences having 50% or higher identity with a majority of the other members of the family: with sequences having identities of 75% or higher are shown in gray. Recombinantly expressed OP/BMP family members which have been shown to make bone are denoted by a "+" in the left margin. In the left margin, TGF-B superfamily members with three-dimensional structures determined are highlighted white on black. The sequences are referenced in Kingsley (Kingsley. (1994) Genes and Development 8:133-146), except for the following: (UNIVIN (Stenzel et al. (1994) Develop. Biol. 166:149-158.), SCREW (Arora, et al. (1994) Genes and Dev. 8:2588-2601.), BMP-9 (Wozney, et al.(1993) PCT/WO 93/00432, SEQ. ID. NO.9), BMP-10 (Celeste et al. (1994) PCT/WO 94/26893, SEQ. ID. NO. 1), GDF-5 (Storm et al. (1994) Nature 368:639-643) (also called CDMP-1 (Chang et al. (1994) J. Biol. Chem. 269: 28227-28234.), GDF-6 (Storm, et al. (1994) Nature 368:639-643), GDF-7 (Storm et al. (1994) Nature 368:639-643), CDMP-2 (Chang et al. (1994) J. Biol. Chem. 269: 28227-28234.), OP-3 (Özkaynak et al. (1994) PCT/WO 94/10203, SEQ. ID. NO. 1), Inhibin Bc (Hötten, et al. (1995) Bioch. Biophys. Res. Comm. 206:608-613), and GDF-10 (Cunningham, et al. (1995) Growth Factors 12:99-109.). The disclosures of the aforementioned citations are incorporated herein by reference. Several sequences in the matrix have alternate names: OP-1 (BMP-7), BMP-2 (BMP-2a), BMP-4 (BMP-2b), BMP-6 (Vgr1), OP-2 (BMP-8), 60A (Vgr-D), BMP-3 (osteogenin), GDF-5 (CDMP-1, MP-52), GDF-6 (CDMP-2, BMP-13) and GDF-7 (CDMP-3, BMP-12).

Figure 7 is a summary of amino acid residues which, according to the 2.8Å resolution structure, together define the solvent accessible surfaces of dimeric hOP-1. Figures 7A, 7B, and 7C show the amino acid sequences defining the human OP-1 finger 1, heel, and finger 2 regions, respectively. The amino acid residues having 40% or greater of their sidechain exposed to solvent are boxed, wherein the solvent accessible amino acid residues that are highly variable among the BMP/OP family of the TGF-B superfamily are identified by shaded boxes.

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Figure 8 is a table, based on the 2.8Å structure, which summarizes the percentage surface accessibility of the amino acid side chains in a hOP-1 monomer subunit and in a hOP-1 dimer. Amino acid residues believed to constitute putative epitopes are designated "EPITOPE" and amino acid residues which are potential candidates as surface modifiable amino acids are marked with an asterisk. In addition, surface modifiable amino acids which are preferred candidates for enhancing solubility are marked with an asterisk.

Figure 9 is a table, based on the 2.8Å structure, which summarizes amino acid residues believed to define the ridge. Amino acid residues believed to constitute the receptor binding domain in the ridge are marked with an asterisk.

Figure 10 is a schematic representation of a computer system useful in the practice of the invention.

Figures 11A and 11B are tables, produced by reference to the 2.8Å structure, which summarize amino acid pairs believed to be useful as sites for introducing additional inter-chain (11A) or intra-chain (11B) disulfide bonds in the hOP-1 dimer.

Figure 12 is an amino acid sequence alignment showing the amino acid sequence of mature human OP-1, and peptides defining the finger-1, finger-2 and heel regions of human OP-1.

Figures 13A-13D are bar graphs illustrating the effect of finger-2 and heel peptides on the alkaline phosphatase activity of ROS cells incubated in either the presence or absence of soluble OP-1. Figures 13A, 13B, 13C, and 13D show the effect of peptides F2-2, F2-3, Hn-2 and Hn-3, respectively, on the alkaline phosphatase activity of ROS cells incubated in the presence (shaded bars) or absence of soluble OP-1 (unshaded bars).

Figures 14A and 14B are graphs showing the displacement of radiolabelled soluble OP-1 from ROS cell membranes by finger 1, finger 2, and heel peptides. Figure 14A shows the displacement of radiolabelled OP-1 from ROS cell membranes by unlabeled soluble OP-1 (open circles and triangles), finger 2 peptide F2-2 (closed circles) and finger 2 peptide F2-3 (closed triangles). Figure 14B shows the displacement of radiolabelled OP-1 from ROS cell membranes by unlabeled soluble OP-1 (open triangles), finger 1 peptide F1-2 (closed boxes), heel peptide H-n2 (open diamonds) and heel peptide H-c2 (open circles).

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Figure 15 is a table summarizing the atomic co-ordinates of hOP-1 resolved to 2.8Å.

Figure 16 is a table summarizing the atomic co-ordinates of hOP-1 resolved to 2.3Å.

Further particulars concerning the drawings are disclosed in the following description which discloses details of the three-dimensional structure of hOP-1, methods for identifying morphogen analogs, and methods for making, testing and using such morphogen analogs.

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#### Detailed Description of Preferred Embodiments

#### 1. Introduction

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As described hereinbelow, the three-dimensional crystal structure of mature hOP-1 now has been solved to 2.3Å. The disclosure provides two sets of atomic co-ordinates for hOP-1, wherein one set of co-ordinates (see Fig. 15) represents the structure of hOP-1 resolved to 2.8Å, and the other set of co-ordinates (see Fig. 16) represents the structure of hOP-1 resolved to 2.3Å. This disclosure thus provides, the atomic co-ordinates defining the relative positions, in three-dimensional space, of at least the C-terminal 104 amino acids of human OP-1 which are sufficient for imparting biological activity. The disclosure provides also an analysis of the structural features of hOP-1. The skilled artisan now can use some or all of these co-ordinates in a database for making morphogenic protein analogs, particularly OP-1 analogs. Specifically, the artisan can select part or all of the database to create templates of part, or all of the hOP-1 structure in three-dimensions, and using this template, create a desired analog or variant which may be amino acid-based, or alternatively composed, in whole or in part, by non-amino acid-based organic components.

Provided below is a detailed description of the three-dimensional crystal structure of hOP-1, along with a detailed description on how to use co-ordinates in a database to design a morphogen analog or structural variant of interest. Amino acid sequences as exemplary templates are provided as examples for designing, identifying, and producing an OP-1 analog using one of the OP-1 atomic co-ordinate databases. Specifically contemplated herein as useful analogs include: small amino molecules which mimic the receptor binding region of the protein; analogs having enhanced stability or solubility; analogs having reduced clearance rates from the body; or enhanced target tissue specificity. The reader will appreciate that these examples are merely exemplary. Given the disclosure of the co-ordinates, the three-dimensional structure, the use of the co-ordinates in a database, and the level of skill in the art today, still other analogs, not specifically recited herein, are contemplated and enabled by this disclosure. In particular, it will be appreciated that, given the disclosure herein, and the known amino acid sequences for other, closely related morphogens, the methods can be used to create other morphogen analogs of, for example, BMP2, BMP4, OP2, BMP5 and BMP6.

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#### II. Structural Determination of hOP-1

#### A. Determination of the 2.8Å Structure

Crystals of mature hOP-1 were grown by mixing equal volumes of purified protein (Özkaynak et al. (1990) EMBO J. 9:2085-20893; and Sampath et al. (1992) J. Biol. Chem. 267:20352-20362) at 10 mg/ml, with 8% saturated ammonium sulfate in 50 mM sodium acetate buffer (pH 5.0) (Griffith et al. (1994) J. Mol. Biol. 244:657-658). The crystals have the symmetry of space group P3<sub>2</sub>21 with unit cell dimensions a=b=99.46Å, and c=42.09Å. One crystal was used to collect a complete native data set to 2.8Å resolution at 4°C. Two heavy atom derivative data sets were collected at 4°C, one from a crystal soaked for seven days in 0.3mM uranyl nitrate and the other from a crystal soaked for eight hours in 0.5 mM sodium gold (III) tetra chloride (Griffith et al. (1994) supra).

The native and derivative data sets were integrated and reduced with the R-AXIS-IIC software suite (Higashi (1990) A Program for Indexing and Processing R-AXIS IIC Imaging Plate Data, Rigaku Corp.) and scaled together using the CCP4 program ANSC (Collaborative Computation Project (1994) Acta Cryst. D50:760-763). Inspection of the Harker sections of the difference Patterson map reveals a single uranyl site. The position of the single gold site was determined by using cross-Fourier techniques using the uranyl position as the phasing site. The heavy atom x,y,z parameters and occupancy were refined with the program TENEYCK (Ten Eyck et al. (1976) J. Mol. Biol. 100:3-11). Using these two derivatives and their anomalous signals, an initial phase set was calculated to 4.0Å resolution with a mean figure of merit of 0.72. The phases were improved and extended to 3.5Å resolution by cycles of solvent flattening (Wang (1985) Meth. Enzymol. 115:90-112) and phase combination (Reed (1986) Acta Cryst. A42:140-149) using the CCP4 (Collaborative Computation Project (1994) supra) crystallographic package. A completely interpretable 3.5Å resolution electron density map permitted the unambiguous tracing of the polypeptide chain and identification of the amino acids from Gln 36 to His 139 using the graphic program "O" (Jones et al. (1991) Acta Crystallogr. A47:110-119). The model was refined with the program XPLOR (Brunger et al. (1987) Science 235:458-460) by using all reflections between 10Å and 2.8Å resolution for which  $F_{obs} > 2.0\sigma$ (F<sub>obs</sub>). There were no water molecules included in the refinement. The root mean square (rms)

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deviation from ideality is 0.02 Å for bond lengths, 3.2° for bond angles. Good stereochemistry was observed for backbone torsion angles. The current R factor is 22.8%.

The atomic co-ordinates defining the 2.8Å resolution structure are listed in Fig. 15. In Fig. 15, the columns entitled "Atom" denote atoms whose co-ordinates have been measured. The first letter in the column defines the element. The columns entitled "Residue" denote the amino acid residues in the hOP-1 monomer which contain an atom whose co-ordinates have been measured. The column entitled "Chain" denotes whether the atom of interest is located within the first (A) or second (B) monomer subunit of the hOP-1 dimer. The columns "X, Y, Z" are the Cartesian co-ordinates that define the atomic position of the atom measured.

#### 10 B. Determination of the 2.3Å Structure

Crystals of mature hOP-1 were produced as described in the previous section. One crystal, frozen in liquid nitrogen, was used to collect a data set to 2.3Å resolution that was 91% complete. The data were collected on imaging plates at beam line X12C (National Synchrotron Light Source) with an oscillation range of 0.5 degrees (overlap of 0.1 degrees) and exposure times of 60-90 seconds. The digitalized data were processed, merged and scaled with DENZO and SCALEPACK (available from Molecular Structure Corporation, Texas). An initial 2Fo-FC map, calculated after X-PLOR rigid-body refinement using the 2.8Å model, was readily interpretable. Portions of the model were manually refitted to the electron-density map with the interactive graphics programs "O" and "Chain". Subsequent cycles of refinement (XPLOR/PROFFT) and manual rebuilding (QUANTA) rapidly converged to the present model.

The current model yielded a conventional crystallographic R factor of 23.5% for data from 10 to 2.3Å (1.5 $\sigma$  cutoff) and a R<sub>free</sub> of 27%. The refined structure was analyzed using the PROCHECK (available from Protein Data Bank, Brookhaven, NY) algorithm and corrected where appropriate. The root mean square (rms) deviation from ideality is 0.015Å for bond distances, 0.034Å for angle distances, and 0.142Å for planar 1-4 distances. The rms deviation from ideality is 1.7 $^{\circ}$  for bond angles. The upper estimate of the error in the atomic positions from the Luzzati plots (EXPLOR) using the free R factor is 0.25-0.33Å. The final model, comprising one monomer subunit, consists of 828 protein atoms (i.e., all non-hydrogen atoms)

and 33 water molecules. The average temperature (B) factor is  $33\text{\AA}^2$  for protein atoms and  $37\text{\AA}^2$  for solvent atoms.

The atomic co-ordinates defining the 2.3Å resolution structure are listed in Figure 16. In Fig. 16, the columns entitled "Atom" denote atoms whose co-ordinates have been measured. The first letter in the column defines the element. The columns entitled "Residue" denote the amino acid residues in the hOP-1 monomer which contain an atom whose co-ordinates have been measured. The column entitled "Chain" denotes whether the atom of interest is located within the first (A) or second (B) monomer subunit of the hOP-1 dimer. The columns "X, Y, Z" are the Cartesian co-ordinates that define the atomic position of the atom measured. The column denoted " $\delta$ " represents the uncertainty in the position of the co-ordinate as derived from the temperature factor (B) of each corresponding atom. The uncertainty of each co-ordinate was derived from the formula  $\delta = \sqrt{\frac{B}{8\pi^2}}$  (see "Protein Crystallography" (1976) T.L. Blundell and L.N. Johnson, Academic Press, p. 121) and is expressed in units of Å.

#### III. Structural Features of hOP-1 Monomer Subunits

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Human OP-1, like TGF-β2, is a dimeric protein having a unique folding pattern involving six of the seven C-terminal cysteine residues, as illustrated in Figure 1A. Each of the subunits in OP-1, like TGF β2 (See Daopin et al. (1992) Science 257:369-373; and Schulnegger et al. (1992) Nature 358:430-434) have a characteristic folding pattern, illustrated schematically in Fig. 1A, that involves six of the seven C-terminal cysteine residues.

Referring to Fig. 1A, four of the cysteine residues in each subunit form two disulfide bonds which together create an eight residue ring, while two additional cysteine residues form a disulfide bond that passes through the ring to form a knot-like structure (cysteine knot). With a numbering scheme beginning with the most N-terminal cysteine of the 7 conserved cysteine residues assigned number 1, the 2nd and 6th cysteine residues are disulfide bonded to close one side of the eight residue ring while the 3rd and 7th cysteine residues are disulfide bonded to close the other side of the ring. The 1st and 5th conserved cysteine residues are disulfide bonded through the center of the ring to form the core of the knot. Amino acid sequence alignment patterns suggest this structural motif is conserved between members of the TGF-β superfamily.

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The 4th cysteine is semi-conserved and when present typically forms an inter-chain disulfide bond (ICDB) with the corresponding cysteine residue in the other subunit.

Each hOP-1 monomer subunit comprises three major tertiary structural elements and an N-terminal region. The structural elements are made up of regions of contiguous polypeptide chain that possess over 50% secondary structure of the following types: (1) loop, (2) helix and (3)  $\beta$ -sheet. Furthermore, in these regions the N-terminal and C-terminal strands are not more than 7 Å apart.

The amino acid sequence between the 1st and 2nd conserved cysteines (Fig. 1A) form a structural region characterized by an anti-parallel  $\beta$ -sheet finger, referred to herein as the finger 1 region (F1). A ribbon trace of the human OP-1 finger 1 peptide backbone is shown in Fig. 1B. Similarly the residues between the 5th and 6th conserved cysteines in Fig. 1A also form an anti-parallel  $\beta$ -sheet finger, referred to herein as the finger 2 region (F2). A ribbon trace of the human OP-1 finger 2 peptide backbone is shown in Fig. 1D. A  $\beta$ -sheet finger is a single amino acid chain, comprising a  $\beta$ -strand that folds back on itself by means of a  $\beta$ -turn or some larger loop so that the entering and exiting strands form one or more anti-parallel  $\beta$ -sheet structures. The third major structural region, involving the residues between the 3rd and 4th conserved cysteines in Fig. 1A, is characterized by a three turn  $\alpha$ -helix referred to herein as the heel region (H). A ribbon trace of the human OP-1 heel peptide backbone is shown in Fig. 1C.

The organization of the monomer structure is similar to that of a left hand (see Fig. 1E) where the knot region is located at the position equivalent to the palm (16), the finger 1 region is equivalent to the index and middle fingers (12 and 13, respectively), the  $\alpha$ -helix, or heel region, is equivalent to the heel of the hand (17), and the finger 2 region is equivalent to the ring and small fingers (14 and 15, respectively). The N-terminal region (undefined in the 2.8 Å resolution map disclosed herein) is predicted to be located at a position roughly equivalent to the thumb (11).

Monovision ribbon tracings illustrating the alpha carbon backbones of each of the three major independent structural elements of the monomer are illustrated in Figures 1B-1D. Specifically, the finger 1 region comprising the first anti-parallel  $\beta$ -sheet segment is shown in

Fig. 1B, the heel region comprising the three turn  $\alpha$ -helical segment is shown in Fig. 1C, and the finger 2 region comprising second and third anti-parallel  $\beta$ -sheet segments is shown in Fig. 1D.

For the sake of comparison, Fig. 3 shows an alignment of the amino acid sequences defining the finger 1, finger 2 and heel regions of hOP-1 and TGF- $\beta$ 2. In Fig. 3, the OP-1 and TGF- $\beta$ 2 amino acid sequences were aligned according to the corresponding regions of local structural identity in the OP-1 and TGF- $\beta$ 2 structures. Alignment gaps were positioned in loop regions, which is where the local conformational homology of the  $\alpha$ -carbon traces tends to be the lowest.

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The structure-based alignment of OP-1 and TGF-β2 then was used as a template for the alignment of the 7-cysteine domain sequences of other TGF-β superfamily members (other members of the TGF-β superfamily are set forth in Fig. 6). Alignment gaps were positioned in regions which are loops in both the OP-1 and TGF-β2 structures. Percent identity between pairs of sequences was calculated as the number of identical aligned sequence positions, excluding gaps, normalized to the geometric mean of the lengths of the sequences and multiplied by 100. Fig. 6 is a matrix of the resulting pair wise present identities between super family sequences so aligned. Using such principles, it is contemplated that the hOP-1 and TGF-β2 structures, either alone or in combination, may be used for homology modeling of other proteins belonging to the TGF-β superfamily whose three-dimensional structures have not yet been determined (see, for example, the other members of the TGF-β superfamily listed in Figure 6). It is contemplated that such models may be useful in designing morphogen analogs for the particular candidate morphogens of interest, however, for simplicity, the disclosure hereinbelow refers specifically the design, identification, and production of morphogen analogs of hOP-1.

Fig. 3 also shows, based on an analysis of the 2.8Å resolution structure, a comparison of interchain contact residues in OP-1 and TGF-β2. Residues were designated as contact residues if the distance between the centers of at least one non-hydrogen atom from each side chain was less than the sum of their Van der Waals radii plus 1.1Å. Despite the low level of sequence identity between OP-1 and TGF-β2, the inter chain contacts between residues in the heel of one chain and residues in finger 1 and finger 2 of the other chain are well conserved.

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Upon detailed inspection of the 2.8Å resolution structure of hOP-1, the finger 1 region of hOP-1 is an antiparallel  $\beta$ -sheet containing a thirteen residue omega loop (Phe 47-Glu 60) (Fig. 2). The structural alignment of the OP-1 and TGF- $\beta$ 2 sequences in Fig. 3 places two gaps in the omega loop. The first gap represents a deletion in hOP-1 that aligns with Arg 26 in the  $\alpha$ 2 helix of TGF- $\beta$ 2. This deletion results in a tighter, non- $\alpha$ -helical turn in OP-1 as compared with TGF- $\beta$ 2. The second gap corresponds to the insertion of Gln 53 in OP-1, which has the result of directing both Gln 53 and Asp 54 side chains into the solvent. By comparison, in the corresponding region of TGF- $\beta$ 2, only Lys 31 is in contact with the solvent. These differences in the conformation of the omega loop also result in the conserved proline (Pro 59) adopting a trans conformation in hOP-1 rather than cis, as in TGF- $\beta$ 2. The conformation of the omega loop orients six non-polar residues so they can contribute to a solvent inaccessible interface with Finger 2. Of these six, four are aromatic (Phe 47, Trp 55, Tyr 62 and Tyr 65), and two are aliphatic (Ile 56 and Ile 57). In all, the conformation of the omega loop backbone places five polar residues (Arg 48, Asp 49, Gln 53, Asp 54, and Glu 60) in contact with solvent. The net surface charge in this region is -2 whereas it is +2 for TGF- $\beta$ 2 (Fig. 5).

According to the 2.8Å structure, the only  $\alpha$  helix in the monomer is located between the third and fifth cysteines (Cys 71 and Cys 104). This helix extends for three and one-half turns from residues Thr 82 to Ile 94, is amphipathic, and contains a number of hydrophobic residues which in the dimer make contact with residues from Finger 1 and Finger 2 of the other monomer (Fig. 3). Several hydrophilic residues (Thr 82, His 84, and Gln 88) form one wall of an internal solvent pocket near the 2-fold axis of the dimer, while others (Asn 83, His 92, and Asn 95) are in contact with the external solvent. The conformation of the loop leading from the C-terminal end of the helix back to the cysteine knot is similar in OP-1 and TGF- $\beta$ 2. By comparison, the loop located at the N-terminal end of the helix is 3 residues longer in OP-1, resulting in a different fold than in TGF- $\beta$ 2. In this loop of OP-1, it is believed that an N-linked sugar moiety is attached to Asn 80, however, no such corresponding glycosylation site exists in TGF- $\beta$ 2. Further, this loop is uncharged in OP-1 whereas it is negatively charged in TGF- $\beta$ 2.

According to the 2.8Å structure, Finger 2 is the second antiparallel β-sheet in OP-1 (Fig. 2). The polypeptide chain reverses direction between segments β6 and β7 through a 3:5 turn (Sibanda, et al. (1991) Methods in Enzymol. 202:59-82) beginning at residue Asp 118 and ending

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at residue Asn 122. In contrast, TGF-β2 has one less residue in this loop and adopts a 2:2 turn (Sibanda et al. (1991) supra). Residues Arg 129 to Val 132, located between segments β7 and β8, form a peptide bridge that crosses over the C-terminal end of strand β5 and produces a 180° twist in the Finger 2 antiparallel β-structure. A similar structure is observed in other cysteine knot growth factors, however the peptide bridge length varies (McDonald et al. (1991) Nature 354:411-414). Within the monomer, Finger 2 makes intra-chain contacts with Finger 1 by contributing aromatic residues Tyr 116, Phe 117 and Tyr 128, and aliphatic residues Val 114, Leu 115, Val 123, Met 131 and Val 133 to a solvent inaccessible interface. OP-1 and TGF-β2 differ by three charges in the region of the Finger 2 turn; OP-1 has two negative charges while TGF-β2 has one positive charge. In the region between the turn and the peptide bridge, OP-1 has a net charge of +3 while TGF-β2 is neutral (Fig. 5).

The N-terminus of each monomeric subunit is believed to be highly mobile and has not been resolved in the 2.8Å resolution structure of hOP-1. The N-terminal region can be deleted without affecting biological activity and, therefore, it is contemplated that this portion of mature hOP-1 may be removed and replaced with other protein or peptide sequences, such as antibodies, and/or radiolabel binding sites for enhancing targeting to a particular locus *in vivo* or for use in *in vivo* imaging experiments. In addition, the N-terminal region may be replaced with an ion chelating motif (e.g., His<sub>6</sub>) for use in affinity purification schemes, or replaced with proteins or peptides for enhancing solubility in aqueous solvents.

#### 20 IV. Structural Features of the hOP-1 Dimer

Fig. 4 shows stereo ribbon trace drawings representative of the peptide backbone of the hOP-1 dimer complex, based on the 2.8Å structure. The two monomer subunits in the dimer complex are oriented symmetrically such that the heel region of one subunit contacts the finger regions of the other subunit with the knot regions of the connected subunits forming the core of the molecule. The 4th cysteine forms an inter-chain disulfide bond with its counterpart on the second chain thereby equivalently linking the chains at the center of the palms. The dimer thus formed is an ellipsoidal (cigar shaped) molecule when viewed from the top looking down the two-fold axis of symmetry between the subunits (Fig. 4A). Viewed from the side, the molecule resembles a bent "cigar" since the two subunits are oriented at a slight angle relative to each other (Fig. 4B).

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As shown in Fig. 4, each of the structural elements which together define the native monomer subunits of the dimer are labeled 43, 43', 44, 44', 45, 45', 46, and 46', wherein, elements 43, 44, 45, and 46 are defined by one subunit and elements 43', 44', 45', and 46' belong to the other subunit. Specifically, 43 and 43' denote the finger 1 regions; 44 and 44' denote heel regions; 45 and 45' denote the finger 2 regions; and 46 and 46' denote disulfide bonds which connect the 1st and 5th conserved cysteines of each subunit to form the knot-like structure. From Fig. 4, it can be seen that the heel region from one subunit, e.g., 44, and the finger 1 and finger 2 regions, e.g., 43' and 45', respectively from the other subunit, interact with one another. These three elements are believed to co-operate with one another to define a structure interactive with the ligand binding interactive surface of the cognate receptor.

The helical axis is defined as the line equi-distant from the alpha carbons in the helical region. A sequence of four points is needed to define the dihedral angle between the axes of the helices in the dimer. The two inner points were chosen to lie on the helical axes adjacent to the  $\alpha$ -carbon of residue His 84 in OP-1 or His 58 in TGF- $\beta$ 2, respectively. The two outer points were chosen to lie on their respective helical axes, but their location is arbitrary. To measure the angle between the helices, the first two points used to define the dihedral angle were translated so as to superimpose the inner points. The resulting three points define the angle.

A major difference between the OP-1 and TGF-β2 dimers is the relative orientation of the helices in the heel region. The angle between the axes of the helices in the heel region of OP-1 is 43° which is 10° larger than that measured for TGF-β2. The measured dihedral angle between the helices is -20° for OP-1 which is 14° more negative than for TGF-β2. Despite these differences in helical orientation, the same helix and finger residue positions are involved in making inter-chain contacts, as evidenced by the shaded residues in Fig. 3.

### A. Differences in the hOP-1 Dimer Relative To Individual Monomer Subunits

During dimerization of the monomer subunits, several amino acids on the surface of each monomer subunit become buried in the hOP-1 dimer. Figure 8 highlights differences in the surface accessibility of particular amino acid residues located in the hOP-1 monomer subunit relative to those in the hOP-1 dimer, as determined from the 2.8 Å structure.

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Loss of non-polar surface area during dimerization was calculated using ACCESS (version 2.1) with a 1.4Å probe (Lee et al. (1971) J. Mol. Biol. 55:379-400). Non-polar surface area is defined as the contribution to the total accessible surface from carbon and sulfur atoms. The surface area measurement algorithm in ACCESS slices the structure into 0.25Å slabs perpendicular to the Z-axis. As a consequence, the results are sensitive to the orientation of a structure relative to the Z-axis (Lee et al. (1971) supra). In order to minimize this effect, we evaluated three perpendicular and one intermediate orientations of each structure. The results of these calculations were combined by accepting, for each non-polar atom, the largest accessible area measured among the four orientations. The values for TGF-β2 reported here were calculated using coordinates from entry 2TG1 (Daopin et al. (1992) supra) and entry 1TFG (Schlunegger et al. (1992) supra) obtained from the January 1994 release of the Protein Data Bank (Bernstein et al. (1977) J. Mol. Biol. 112:535-542) at Brookhaven National Laboratory.

In Figure 8, the column entitled "Residue" denotes an amino acid of interest. The column entitled "Monomer % Area" denotes the percentage of the amino acid that is exposed on the surface of the hOP-1 monomer, the column entitled "Dimer % Area" denotes the percentage of the amino acid that is exposed on the surface of the hOP-1 dimer, and the column denoted "Hidden % Area" denotes amount of surface area for each amino acid that is lost upon dimerization of each monomer subunit to produce the hOP-1 dimer. This analysis reveals amino acids which become buried during dimerization and, thus, likely are located at the interface of the two monomer subunits. For example, 70.75% of the surface area of His 84 becomes hidden upon dimerization. A review of the structure of dimeric hOP-1 reveals that His 84 is located at the interface between the two monomers.

#### B. Solution Electrostatic Potentials on the Surface of OP-1 and TGF-B2

The solution electrostatic potentials surrounding the OP-1 and TGF-β2 (1TFG)

(Schlunegger et al. (1992) supra) dimers were calculated using DELPHI (Gilson et al. (1987)

Nature 330:84-86; and Nicholls et al. (1991) J. Comput. Chem. 12:435-445) (Biosym

Technologies, Inc., San Diego, CA). The calculations were performed using a solvent dielectric constant of 80, a solvent radius of 1.4Å, an ionic strength of 0.145M and an ionic radius of 2.0 Å. The interior of the protein was modeled using a dielectric constant of 2.0. Formal charges were used and distributed as follows: atoms OD1 and OD2 of Asp were each charged -0. 5, atoms

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OE1 and OE2 of Glu were each charged -0.5, atoms ND1 and NE2 of His were each charged 0.25, atom NZ of Lys was charged +1.0, atoms NH1 and NH2 of Arg were each charged +0.5, and atom OXT of the C-terminal carboxyl group was charged -1.0.

The differences in charge distribution on the surfaces of OP-1 and TGF-\(\beta\)2 can be observed by comparing the color distributions of Figs. 5B and 5C, respectively. Surface regions having an electrostatic potential of -3kT or less are shown in red while surface regions of +3kT or greater are shown in blue. Neutral regions are shown in green or gold to correspond to the backbone ribbons shown in Fig. 5A. As mentioned in the following section, the differences in electrostatic potential on the surfaces of OP-1 and TGF-\(\beta\)2 may play an important role in the specific interactions of the TGF-\(\beta\) superfamily members with their cognate receptors.

#### C. Receptor Binding Domain

Without wishing to be bound by theory, it is contemplated that the receptor binding regions of hOP-1 includes amino acids that are both solvent accessible and lie at positions of heterogeneous composition, as determined from the amino acid sequence of hOP-1 when aligned with other members of the TGF- $\beta$  superfamily (See Fig. 3). Divergent structural features in hOP-1, like TGF- $\beta$ 2, occur primarily in the external loops of finger 1 and finger 2, the loops bordering the helix in the heel region, and the residues in the N-terminal domain preceding the first cysteine of the cysteine knot. These regions are solvent accessible. In both the OP-1 and TGF- $\beta$ 2 dimer structures, the tip of finger 2 and the omega loop of finger 1 from one chain, and the C-terminal end of the  $\alpha$ -helix in the heel of the other chain form a contiguous ridge approximately 40 Å long and 15 Å wide (Fig. 5A). It is contemplated that this ridge contains the primary structural features that interact with the cognate receptor, and that the binding specificity between different TGF- $\beta$  superfamily members derives from conformational and electrostatic variations on the surface of this ridge.

Differences in the conformation of the finger 1 omega loop, which constitutes the mid section of the ridge, and in the turn at the end of finger 2, which forms one end of the ridge are noted. However, there are striking differences in the surface charge of the ridge in hOP-1 relative to TGF-\beta2 (see Figs. 5B and 5C). In hOP-1, the ends of the finger regions are negatively charged whereas in TGF-\beta2, the ends of the finger regions are positively charged. This results in

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a net charge of -4 for the receptor binding ridge of hOP-1 versus +3 for TGF- $\beta$ 2. Conversely, the  $\beta$ -strand located C-terminal to the turn of finger 2 ( $\beta$ 7, Fig. 2) is positively charged in OP-1 whereas it is negatively charged in TGF- $\beta$ 2 (Figs. 5B and 5C). These features suggests that electrostatic charge distribution plays an important role in the specific interactions of the TGF- $\beta$  superfamily members with their cognate receptors.

Figure 9 summarizes the amino acid residues which, according to the 2.8 Å structure, are believed to constitute the ridge, and also indicates whether each amino acid residue is disposed within the heel, finger 1, or finger 2 domains. Figure 9 also provides a list of amino acid residues which are believed to constitute at least part, if not all of the receptor binding domain of hOP-1.

#### V. Design of Morphogen Analogs

Although it is contemplated that the design of morphogen analogs can be facilitated by conventional ball and stick type modeling procedures, it is contemplated that the ability to design morphogen analogs is enhanced significantly using modern computer-driven modeling and design procedures.

It is contemplated that the design of morphogen analogs, as discussed in detail hereinbelow, is facilitated using conventional molecular modeling computers or workstations, commercially available from, for example, Silicon Graphics, Inc. or Evans and Sutherland Computer Corp., which implement equally conventional computer modeling programs, for example, INSIGHTII, DISCOVER, and DELPHI, commercially available from Biosym, Technologies Inc., and QUANTA, and CHARMM commercially available from Molecular Simulations, Inc.

Furthermore, it is understood that any computer system having the overall characteristics set forth in Fig. 10 may be useful in the practice of the instant invention. More specifically, Fig. 10, is a schematic representation of a typical computer work station having in electrical communication (100) with one another via, for example, an internal bus or external network, a processor (101), a RAM (102), a ROM (103), a terminal (104), and optionally an external storage device, for example, a diskette, CD ROM, or magnetic tape (105).

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It is contemplated, that the co-ordinates can be used not only to provide a basis for reengineering hOP-1 dimers by using, for example, site-directed mutagenesis methodologies, to enhance, for example, the solubility and or/stability of the active hOP-1 dimer in physiological buffers, but also to provide a starting point for the *de novo* design and production of peptides or other small molecules which mimic the bioactivity of hOP-1. Set forth below are illustrative examples demonstrating the usefulness of hOP-1 atomic co-ordinates in the design of morphogen analogs, however, it is understood the examples below are illustrative and not meant to be limiting in any way.

#### A. Engineering hOP-1 Dimers

In one aspect, the availability of the atomic co-ordinates for hOP-1, enables the artisan to perform theoretical amino acid replacements and to determine by calculation, in advance of actually making and testing the candidate molecule in a laboratory setting, whether a particular amino acid substitution disrupts the packing of the OP-1 dimer and whether a morphogen analog is likely to be more stable and/or soluble than the template OP-1 molecule. Such procedures assist the artisan to eliminate non viable replacements and to focus efforts on more promising candidate analogs.

#### (i) Enhancing the Stability of hOP-1 Dimers

It is contemplated that the skilled artisan in possession of the atomic co-ordinates defining hOP-1 can introduce additional inter- or intra-chain covalent and/or non-covalent interactions into the hOP-1 dimer to stabilize the dimer by preventing disassociation or unfolding of each monomer subunit. Preferred engineered covalent interactions include, for example, engineered disulfide bonds, and preferred engineered non-covalent interactions include, for example, hydrogen bonds, salt bridges, and hydrophobic interactions.

For example, in order to introduce additional disulfide bonds, the skilled artisan can identify sites suitable for the introduction of a pair of cysteine amino acid residues by using standard molecular modeling programs, for example, INSIGHT, DISCOVER, CHARMM and QUANTA. Another program useful in identifying pairs of amino acids as potential sites for introducing stabilizing disulfide bonds is described in U.S. Patent No. 4,908,773, the disclosure of which is incorporated herein by reference.

For example, the skilled artisan using the INSIGHT program can screen for pairs of amino acids, where the distance between the C $\beta$  atoms of each amino acid is in the range of about 3.0 to about 5.0, or more preferably about 3.5 to about 4.5 Å apart. For this purpose, glycines, which contain no C $\beta$ -C $\beta$  bond, are first converted to alanines on the computer. The possible range of C $\beta$ -C $\beta$  distances in a disulfide bond are 3.1 Å to 4.6 Å, but separations outside this range can be accommodated by small shifts in the neighboring atoms. Searching C $\beta$ , rather than C $\alpha$  distances, ensures both reasonable spacing as well as proper orientation of the C $\alpha$ -C $\beta$  bond. The effects of adding such an additional linkage on protein structure are determined by mutating the two candidates residues to Cys; rotating each new Cys about the C $\alpha$ -C $\beta$  bond to bring the two  $\gamma$  sulfurs as close to within 2Å as possible; creating a disulfide between the  $\gamma$  sulfurs; and energy minimizing structural regions within 5 Å of the disulfide bond. Any deformation of the structure caused by introduction of the additional disulfide bond is revealed by inspection when the minimized, mutated model structure is superimposed on the native structure.

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It is contemplated that the introduction of additional linkages will improve solubility by preventing transient exposure of non-polar interface or buried residues. Figure 11A lists amino acid residues, based on the 2.8Å structure, which may be mutated to cysteine residues for introducing additional inter-chain disulfide bonds, based upon the selection criteria presented above. For reference purposes, Table 11A includes also the length of the naturally occurring inter-chain disulfide linkage in wild type hOP-1, that is, the disulfide linkage connecting Cys-103 of one monomer subunit with the counterpart Cys-103 of the other monomer subunit.

A preferred pair of residues suitable for modification include the residue at position 83 of one chain and the residue at position 130 of the other chain. It is contemplated that the additional inter-chain linkage stabilizes the dimeric structure by connecting the N-terminal end of the Heel helix of the first subunit to the middle of the Finger 2 region in the second subunit. A disulfide bond between position 82 on one chain and position 130 of the other chain also is geometrically feasible, but because Thr 82 is part of the NAT glycosylation site in OP-1, its modification may inhibit proper glycosylation.

Figure 11B summarizes amino acid residues which can be mutated to cysteine residues for introducing additional intra-chain disulfide bonds, based upon the selection criteria presented

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above. As noted previously, the putative receptor binding region comprises at least two physically proximal, but sequentially separate regions, namely the tips of Finger 1 and Finger 2. It is contemplated that the structural integrity of the putative receptor binding ridge can be stabilized by engineering an intra-chain disulfide link between residues of Finger 1 and Finger 2. In a preferred embodiment, the residue at position 58 in Finger 1 can be disulfide bonded with the residue at position 114 in Finger 2. It is contemplated that a link between the residues at positions 58 and 115 also would be viable, however, this would move the disulfide bond nearer to the putative receptor binding region on Finger 2. Also a link between positions 65 and 133 is possible, however, this would be located near to the knot region of each chain and, thus may have little effect on stabilizing the putative receptor binding regions at the tips of Finger 1 and Finger 2. Additionally, the proximity of such a linkage to the disulfide bonds in the knot region might interfere with the proper formation of those structures.

With regard to non-covalent interactions, it is contemplated that the structural stability of the hOP-1 dimer can be enhanced by increasing inter-chain hydrogen bonding.

The electrostatic potential (due to other charges in a protein) in the region of a charged residue affects the pK of that residue. Because the pK's of both histidine and the N-terminal primary amino group are near neutrality, it may be possible to modify their pKs through the placement of charges on the surface of the molecule. It is contemplated that the buried His at position His 84 in hOP-1 helps stabilize the structure of the dimer by participating in hydrogen bonds with backbone carboxyl groups of residues Ala 64 and Tyr 65 of the other chain.

Accordingly, it is contemplated that the introduction of surface charges may enhance this effect and thereby further stabilize the structure of the molecule. For example, mutating Tyr 65 or Val 132 to Asp may further polarize the carbonyl bonds of the amino acid residues at positions 64 and 65, as well as raise the pK of His 84. The pK of His 84 may further be affected by replacing residues Tyr 44, Ala 63, or Asn 110 by an Asp. It is contemplated that the preferred modification for this purpose is Tyr 65-> Asp 65.

Using the same basic principles, the skilled artisan likewise can identify pairs of amino acids whose replacement can facilitate the introduction of an inter-chain salt bridge, internal hydrogen bond, or hydrophobic interaction. Such determinations are within the scope of an artisan having an ordinary level of skill in the field of molecular modeling.

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Once a pair of target amino acids has been identified, the site-directed replacement of the target amino acids with the desirable replacements can be facilitated by the use of conventional site-directed mutagenesis procedures, for example, by cassette mutagenesis or oligonucleotide-directed mutagenesis. Such techniques are thoroughly documented in the art and so are not discussed herein. The effect of the site-specific replacements on the stability of resulting modified hOP-1 dimers or muteins can be measured, after production and purification, using standard methodologies well known in the art, for example, circular dichroism, analytical centrifugation, differential scanning calorimetry, fluoresence or other spectroscopic techniques.

#### (ii) Enhancing Water Solubility of hOP-1 Dimer

OP-1 has limited solubility in aqueous solvents. It is contemplated, however, that by using the hOP-1 atomic co-ordinates that the artisan can replace amino acids at the solvent accessible surface of the dimer thereby to increase the dielectric properties of dimeric hOP-1. For example, solvent accessible hydrophobic amino acid residues, such as, glycine, alanine, valine, leucine and isoleucine may be replaced by more polar residues, such as, lysine, arginine, histidine, aspartate, asparagine, glutamate and glutamine.

The solvent accessible amino acids can be identified using a computer program, such as ACCESS (version 2.1) using a 1.4 Å probe (Lee et al. (1972) supra). In Fig. 7 amino acid residues having at least 20% of their side chain areas exposed to solvent are boxed. When modifying surface residues it is important not to produce new epitopes that can be recognized as non-host especially, if the hOP-1 analogs are to be used as injectable molecules. It is believed that, amino acid side chains seen by a 10 Å spherical probe likely are part of surface epitopes. One skilled in the art can use ACCESS with a 10 Å spherical probe to identify potential epitopes, however this process can be carried out manually using a graphics package, such as, INSIGHT II. In Figure 8, residue side chains so identified as potential epitopes are highlighted. Residue positions that are candidates for modification so as to improve the solubility of the dimer are highlighted. Preferred candidate amino acids for replacement include, for example, Ala 63, Ala 72, Ala 81, Ala 111 and Ala 135, Ile 86, Ile 112, Tyr 52, Tyr 65, and Tyr 128.

Once solvent accessible hydrophobic or non polar amino acids have been identified (see Fig. 9), these amino acids theoretically may be replaced, via a computer, with more polar amino

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acids. The effect of the amino acid replacements on the solution electrostatic potentials surrounding the modified hOP-1 dimer as well as the free energy of the dimer can calculated using the program DELPHI (Gilson et al (1987) supra; Nicholls et al. (1991) supra). Preferred amino acid substitutions lower the free energy of the hOP-1 dimer without introducing potential antigenic sites. As mentioned above, such antigenic sites may be detected by implementing a computer program like ACCESS (version 2.1) using a 10 Å probe. In addition, it is contemplated that preferred surface residues suitable for replacement do not constitute part of the receptor binding domain.

The resulting candidate morphogen analogs can be produced using conventional site-directed mutagenesis methodologies and the effect of the site-directed modification on the solubility of the hOP-1 dimer can be measured, for example, by comparing the partition coefficient or "salting out properties" of the modified hOP-1 dimer versus the native hOP-1 dimer. See for example, Scopes (1987) in *Protein Purification: Principles and Practice, 2nd Edition* (Springer-Verlag); and Englard et al. (1990) Methods in Enzymology 182: 285-300.

#### (iii) Engineering Glycosylation Sites

In addition to replacing single, solvent accessible amino acid residues with more polar or hydrophobic amino acid residue, one or more solvent accessible amino acid residues may be replaced so as to create a new eukaryotic glycosylation site or alternatively to eliminate or alter an existing glycosylation site. Glycosylation sites are well known and are thoroughly described in the art. Addition of a new glycosylation site or alteration of an existing site may result in the addition of one or more glycosyl groups, e.g., N-acetyl-sialic acid, which may enhance the solubility of the morphogen analog. As described herein, such sites can be introduced by site-directed mutagenesis methodologies which are well known in the art. Preferably, such sites do not create new antigenic determinants (although these may be tolerable for short duration therapeutic uses). Reference to Table 8 identifies surface accessible amino acid residues, based on the 2.8Å structure, which likely are not part of an antigenic epitope and which may be used as candidates for introducing an additional glycosylation site.

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### B. Engineering Small Molecules Based Upon The hOP-1 Structure

The availability of atomic co-ordinates for hOP-1 enables the skilled artisan to design small molecules, for example, peptides or non-peptidyl based organic molecules having certain chemical features, which mimic the biological activity of hOP-1. Chemical features of interest may include, for example, the three-dimensional structure of a particular protein domain, solvent accessible surface of a particular protein domain, spatial distribution of charged and/or hydrophobic chemical moieties, electrostatic charge distribution, or a combination thereof. Such chemical features may readily be determined from the three-dimensional representation of hOP-1.

#### (i) Peptides

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After having determined which amino acid residues contribute to the receptor binding domain (*supra*), it is possible for the skilled artisan to design synthetic peptides having amino acid sequences that define a pre-selected receptor binding motif. A computer program useful in designing potentially bioactive peptido-mimetics is described in U.S. Patent No. 5,331,573, the disclosure of which is incorporated by reference herein.

In addition to choosing a desirable amino acid sequence, the skilled artisan using standard molecular modeling software packages, *infra*, can design specific peptides having, for example, additional cysteine amino acids located at pre-selected positions to facilitate cyclization of the peptide of interest. Oxidation of the additional cysteine residues results in cyclization of the peptide thereby constraining the peptide in a conformation which mimics the conformation of the corresponding amino acid sequence in native hOP-1. It is contemplated, that any standard covalent linkage, for example, disulfide bonds, typically used to cyclize synthetic peptides maybe useful in the practice of the instant invention. Alternative cyclization chemistries are discussed in International Application PCT/WO 95/01800, the disclosure of which is incorporated herein by reference.

In addition, it is contemplated that a single peptide containing amino acid sequences derived from separate hOP-1 subunit domains, for example, a single peptide having an amino acid sequence defining the tip of the finger 1 region linked by means of a polypeptide linker to an amino acid sequence defining the tip of the finger 2 region. The amino sequence defining

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each of the finger regions may further comprise a means, for example, disulfide bonds for cyclizing each finger region motif. The resulting peptide therefore comprises a single polypeptide chain having a first amino acid sequence defining a three-dimensional domain mimicking the tip of the finger 1 region and a second said sequence defining a three-dimensional domain mimicking the tip of the finger 2 region.

Such peptides may be synthesized and screened for OP-1 like activity using any of the standard protocols described below.

#### (ii) Organic molecules

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As discussed above, upon determination of the receptor binding domain of hOP-1, it is contemplated that the skilled artisan, can design non-peptidyl based small molecules, for example, small organic molecules, whose structural and chemical features mimic the same features displayed on at least part of the surface of the receptor binding domain of hOP-1.

Because a major contribution to the receptor binding surface is the spatial arrangement of chemically interactive moieties present within the sidechains of amino acids which together define the receptor binding surface, a preferred embodiment of the present invention relates to designing and producing a synthetic organic molecule having a framework that carries chemically interactive moieties in a spatial relationship that mimics the spatial relationship of the chemical moieties disposed on the amino acid sidechains which constitute the receptor binding site of hOP-1. Preferred chemical moieties, include but are not limited to, the chemical moieties defined by the amino acid side chains of amino acids believed to constitute the receptor binding domain of hOP-1 (See Fig. 9). It is understood, therefore, that the receptor binding surface of the morphogen analog need not comprise amino acid residues but the chemical moieties disposed thereon.

For example, upon identification of relevant chemical groups, the skilled artisan using a conventional computer program can design a small molecule having the receptor interactive chemical moieties disposed upon a suitable carrier framework. Useful computer programs are described in, for example, Dixon (1992) *Tibtech 10*: 357-363; Tschinke et al. (1993) *J. Med. Chem 36*: 3863-3870; and Eisen et al. (1994) *Proteins: Structure, Function, and Genetics 19*: 199-221, the disclosures of which are incorporated herein by reference.

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One particular computer program entitled "CAVEAT" searches a database, for example, the Cambridge Structural Database, for structures which have desired spatial orientations of chemical moieties (Bartlett *et al.* (1989) in "Molecular Recognition: Chemical and Biological Problems" (Roberts, S.M., ed) pp 182-196). The CAVEAT program has been used to design analogs of tendamistat, a 74 residue inhibitor of  $\alpha$ -amylase, based on the orientation of selected amino acid side chains in the three-dimensional structure of tendamistat (Bartlett *et al.* (1989) supra).

Alternatively, upon identification of a series of analogs which mimic the biological activity of OP-1, as determined by *in vivo* or *in vitro* assays, the skilled artisan may use a variety of computer programs which assist the skilled artisan to develop quantitative structure activity relationships (QSAR) and further to assist in the *de novo* design of additional morphogen analogs. Other useful computer programs are described in, for example, Connolly-Martin (1991) *Methods in Enzymology 203:*587-613; Dixon (1992) *supra*; and Waszkowycz *et al.* (1994) *J. Med. Chem. 37*: 3994-4002.

Thus, for example, one can begin with a portion of the three dimensional structure of OP-1 (or a related morphogen) corresponding to a region of known or suspected biological importance. One such region is the solvent accessible loop or "tip" of the finger 2 region between the \$6 and \$7 sheets (i.e., from approximately residues 118-122). Synthetic, cyclic peptides (i.e., F2-2 and F2-3) were produced including this region (and several flanking residues) and were shown to possess OP-1-like biological activity (see Examples below). Based upon the three-dimensional structure of this region, disclosed herein, one is now enabled to produce more effective OP-1-like (or, generally, morphogen-like) analogs. For example, as shown in great detail in Figures 7-9 and 15, the charged γ-carboxy groups of Asp 118 and Asp 119, and the relatively hydrophilic hydroxyl groups of Ser 120 and Ser 121, are solvent accessible and believed to be involved in OP-1 receptor binding. The relative positions of these groups in three dimensions in OP-1 are given in Figures 15 and 16. These functional groups define a contiguous portion of the three dimensional structure of the OP-1 surface. The peptide backbone of these residues, however, is not solvent accessible and, therefore, is not believed to form a portion of the three-dimensional surface of the OP-1 molecule. Thus, one of ordinary skill in the art, when choosing or designing an OP-1 or morphogen analog, can choose or design a molecule having

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the same or substantially equivalent (e.g., thiol v. hydroxyl) functional groups in substantially the same (e.g.,  $\pm$  1-3 Å) three-dimensional conformation. The same is true for other regions of interest in the OP-1 monomers or dimers (e.g., the receptor binding domain, the finger 1, finger 2, or heel regions, or solvent accessible portions thereof). By using the three-dimensional structures disclosed herein, including the disclosure of the positions of solvent accessible and probable receptor contact residues, one of ordinary skill in the art can choose a portion of the three-dimensional structure of the OP-1 (or a related morphogen) molecule and, using this "portion" as a template select or design an analog which functionally mimics the template structure.

The molecular framework or backbone of the morphogen analog can be freely chosen by one of ordinary skill in the art so that it (1) joins the functional groups which mimic the portion of the morphogen's contiguous three-dimensional surface, including charge distribution and hydrophobicity/hydrophilicity characteristics, and (2) maintains or, at least, allows the functional groups to maintain the appropriate three-dimensional surface interaction and spatial relationships, including any hydrogen bonding and electrostatic interactions. As described above, peptides are obvious choices for the production of such morphogen analogs because they can provide all of the necessary functional groups and can assume appropriate three-dimensional structures. Several examples of peptide analogs of the finger regions are described herein, below. The peptides are cyclized to maintain hydrogen bonds and create a structure which mimics that of the template. These peptides are synthesized from a linear primary sequence of amino acids in finger 2. An alternative peptide can be created, for example, which combines portions of finger 1 and finger 2, constructed to mimic the structure of the tips of fingers 1 and 2 together as they occur in the folded OP1 monomer. Biologically active peptides such as F2, F3 or others, then can be used as is or, more preferably, become lead compounds for iterative modification to create a compound that is more stable or more active in vivo. For example, the peptide backbone can be reduced or replaced to reduce hydrolysis in vivo. Alternatively, structural modifications can be introduced to the backbone or by amino acid substitutions which more accurately mimic the protein's structure when bound to the receptor. These second generation structures then can be tested for enhanced binding. In addition, iterative amino acid

replacements with alanines, ("alanine scan") can be used to determine the minimum residue contacts required for binding.

Once these minimum functional groups are known, a fully synthetic molecule can be created which mimics the charge or electrostatic distribution of the minimum required functional groups, and provides the appropriate bulk and structure to functionally mimic a second generation molecule having the desired binding affinity.

#### VI. Production of Morphogen Analogs.

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As mentioned above, the morphogen analogs of the invention may comprise modified hOP-1 dimeric proteins or small molecules, for example, peptides or small organic molecules. It is contemplated that any appropriate methods can be used for producing a pre-selected morphogen analog. For example, such methods may include, but are not limited to, methods of biological production from suitable host cells or synthetic production using synthetic organic chemistries.

For example, modified hOP-1 dimeric proteins or hOP-based peptides may be produced using conventional recombinant DNA technologies, well known and thoroughly documented in the art. Under these circumstances, the proteins or peptides may be produced by the preparation of nucleic acid sequences encoding the respective protein or peptide sequences, after which, the resulting nucleic acid can be expressed in an appropriate host cell. By way of example, the proteins and peptides may be manufactured by the assembly of synthetic nucleotide sequences and/or joining DNA restriction fragments to produce a synthetic DNA molecule. The DNA molecules then are ligated into an expression vehicle, for example an expression plasmid, and transfected into an appropriate host cell, for example *E. coli*. The protein encoded by the DNA molecule then is expressed, purified, folded if necessary, tested *in vitro* for binding activity with an OP-1 receptor, and subsequently tested to assess whether the morphogen analog induces or stimulates hOP-1-like biological activity.

The processes for manipulating, amplifying, and recombining DNA which encode amino acid sequences of interest generally are well known in the art, and therefore, are not described in detail herein. Methods of identifying and isolating genes encoding hOP-1 and its cognate receptors also are well understood, and are described in the patent and other literature.

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Briefly, the construction of DNAs encoding the biosynthetic constructs disclosed herein is performed using known techniques involving the use of various restriction enzymes which make sequence specific cuts in DNA to produce blunt ends or cohesive ends, DNA ligases, techniques enabling enzymatic addition of sticky ends to blunt-ended DNA, construction of synthetic DNAs by assembly of short or medium length oligonucleotides, cDNA synthesis techniques, polymerase chain reaction (PCR) techniques for amplifying appropriate nucleic acid sequences from libraries, and synthetic probes for isolating OP-1 genes or genes encoding other members of the TGF- $\beta$  superfamily as well as their cognate receptors. Various promoter sequences from bacteria, mammals, or insects to name a few, and other regulatory DNA sequences used in achieving expression, and various types of host cells are also known and available. Conventional transfection techniques, and equally conventional techniques for cloning and subcloning DNA are useful in the practice of this invention and known to those skilled in the art. Various types of vectors may be used such as plasmids and viruses including animal viruses and bacteriophages. The vectors may exploit various marker genes which impart to a successfully transfected cell a detectable phenotypic property that can be used to identify which of a family of clones has successfully incorporated the recombinant DNA of the vector.

One method for obtaining DNA encoding the biosynthetic constructs disclosed herein is by assembly of synthetic oligonucleotides produced in a conventional, automated, oligonucleotide synthesizer followed by ligation with appropriate ligases. For example, overlapping, complementary DNA fragments may be synthesized using phosphoramidite chemistry, with end segments left unphosphorylated to prevent polymerization during ligation. One end of the synthetic DNA is left with a "sticky end" corresponding to the site of action of a particular restriction endonuclease, and the other end is left with an end corresponding to the site of action of another restriction endonuclease. The complementary DNA fragments are ligated together to produce a synthetic DNA construct.

After the appropriate DNA molecule has been synthesized, it may be integrated into an expression vector and transfected into an appropriate host cell for protein expression. Useful prokaryotic host cells include, but are not limited to, *E. coli*, and *B. subtilis*. Useful eukaryotic host cells include, but are not limited to, yeast cells, insect cells, myeloma cells, fibroblast 3T3 cells, monkey kidney or COS cells, chinese hamster ovary (CHO) cells, mink-lung epithelial

cells, human foreskin fibroblast cells, human glioblastoma cells, and teratocarcinoma cells.

Alternatively, the genes may be expressed in a cell-free system such as the rabbit reticulocyte lysate system.

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The vector additionally may include various sequences to promote correct expression of the recombinant protein, including transcriptional promoter and termination sequences, enhancer sequences, preferred ribosome binding site sequences, preferred mRNA leader sequences, preferred protein processing sequences, preferred signal sequences for protein secretion, and the like. The DNA sequence encoding the gene of interest also may be manipulated to remove potentially inhibiting sequences or to minimize unwanted secondary structure formation. The morphogenic protein analogs proteins also may be expressed as fusion proteins. After being translated, the protein may be purified from the cells themselves or recovered from the culture medium and then cleaved at a specific protease site if so desired.

For example, if the gene is to be expressed in *E. coli*, it is cloned into an appropriate expression vector. This can be accomplished by positioning the engineered gene downstream of a promoter sequence such as Trp or Tac, and/or a gene coding for a leader peptide such as fragment B of protein A (FB). During expression, the resulting fusion proteins accumulate in refractile bodies in the cytoplasm of the cells, and may be harvested after disruption of the cells by French press or sonication. The isolated refractile bodies then are solubilized, and the expressed proteins folded and the leader sequence cleaved, if necessary, by methods already established with many other recombinant proteins.

Expression of the engineered genes in eukaryotic cells requires cells and cell lines that are easy to transfect, are capable of stably maintaining foreign DNA with an unrearranged sequence, and which have the necessary cellular components for efficient transcription, translation, post-translation modification, and secretion of the protein. In addition, a suitable vector carrying the gene of interest also is necessary. DNA vector design for transfection into mammalian cells should include appropriate sequences to promote expression of the gene of interest as described herein, including appropriate transcription initiation, termination, and enhancer sequences, as well as sequences that enhance translation efficiency, such as the Kozak consensus sequence. Preferred DNA vectors also include a marker gene and means for amplifying the copy number of the gene of interest. A detailed review of the state of the art of the production of foreign proteins

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in mammalian cells, including useful cells, protein expression-promoting sequences, marker genes, and gene amplification methods, is disclosed in Bendig (1988) *Genetic Engineering* 7:91-127.

The best characterized transcription promoters useful for expressing a foreign gene in a particular mammalian cell are the SV40 early promoter, the adenovirus promoter (AdMLP), the mouse metallothionein-I promoter (mMT-I), the Rous sarcoma virus (RSV) long terminal repeat (LTR), the mouse mammary tumor virus long terminal repeat (MMTV-LTR), and the human cytomegalovirus major intermediate-early promoter (hCMV). The DNA sequences for all of these promoters are known in the art and are available commercially.

The use of a selectable DHFR gene in a dhfr- cell line is a well characterized method useful in the amplification of genes in mammalian cell systems. Briefly, the DHFR gene is provided on the vector carrying the gene of interest, and addition of increasing concentrations of the cytotoxic drug methotrexate, which is metabolized by DHFR, leads to amplification of the DHFR gene copy number, as well as that of the associated gene of interest. DHFR as a selectable, amplifiable marker gene in transfected chinese hamster ovary cell lines (CHO cells) is particularly well characterized in the art. Other useful amplifiable marker genes include the adenosine deaminase (ADA) and glutamine synthetase (GS) genes.

The choice of cells/cell lines is also important and depends on the needs of the experimenter. COS cells provide high levels of transient gene expression, providing a useful means for rapidly screening the biosynthetic constructs of the invention. COS cells typically are transfected with a simian virus 40 (SV40) vector carrying the gene of interest. The transfected COS cells eventually die, thus preventing the long term production of the desired protein product but provide a useful technique for testing preliminary analogs for binding activity.

The various cells, cell lines and DNA sequences that can be used for mammalian cell expression of the single-chain constructs of the invention are well characterized in the art and are readily available. Other promoters, selectable markers, gene amplification methods and cells also may be used to express the proteins of this invention. Particular details of the transfection, expression, and purification of recombinant proteins are well documented in the art and are understood by those having ordinary skill in the art. Further details on the various technical

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aspects of each of the steps used in recombinant production of foreign genes in mammalian cell expression systems can be found in a number of texts and laboratory manuals in the art, such as, for example, Ausubel et al., ed., Current Protocols in Molecular Biology, John Wiley & Sons, New York, (1989).

Alternatively, morphogen analogs which are small peptides, usually up to 50 amino acids in length, may be synthesized using standard solid-phase peptide synthesis procedures, for example, procedures similar to those described in Merrifield (1963) *J. Am. Chem. Soc.*, 85:2149. For example, during synthesis, N-α-protected amino acids having protected side chains are added stepwise to a growing polypeptide chain linked by its C-terminal end to an insoluble polymeric support, e.g., polystyrene beads. The peptides are synthesized by linking an amino group of an N-α-deprotected amino acid to an α-carboxy group of an N-α-protected amino acid that has been activated by reacting it with a reagent such as dicyclohexylcarbodiimide. The attachment of a free amino group to the activated carboxyl leads to peptide bond formation. The most commonly used N-α-protecting groups include Boc which is acid labile and Fmoc which is base labile.

Briefly, the C-terminal N-α-protected amino acid is first attached to the polystyrene beads. Then, the N-α-protecting group is removed. The deprotected α-amino group is coupled to the activated α-carboxylate group of the next N-α-protected amino acid. The process is repeated until the desired peptide is synthesized. The resulting peptides are cleaved from the insoluble polymer support and the amino acid side chains deprotected. Longer peptides, for example greater than about 50 amino acids in length, typically are derived by condensation of protected peptide fragments. Details of appropriate chemistries, resins, protecting groups, protected amino acids and reagents are well known in the art and so are not discussed in detail herein. See for example, Atherton et al. (1963) Solid Phase Peptide Synthesis: A Practical Approach (IRL Press,), and Bodanszky (1993) Peptide Chemistry, A Practical Textbook, 2nd Ed, Springer-Verlag, and Fields et al. (1990) Int. J. Peptide Protein Res. 35:161-214, the disclosures of which are incorporated herein by reference.

Purification of the resulting peptide is accomplished using conventional procedures, such as preparative HPLC, e.g., gel permeation, partition and/or ion exchange chromatography. The

choice of appropriate matrices and buffers are well known in the art and so are not described in detail herein.

With regard to the production of non-peptide small organic molecules which induce OP-1 like biological activities, these molecules can be synthesized using standard organic chemistries well known and thoroughly documented in the patent and other literatures.

## VII. Screening For Binding and Biological Activity.

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As a first step in determining whether a morphogen analog induces an OP-1 like biological activity, the skilled artisan can use a standard ligand-receptor assay to determine whether the morphogen analog binds preferentially to OP-1 receptor. For standard receptor-ligand assays, the artisan is referred to, for example, Legerski et al. (1992) Biochem. Biophys. Res. Comm. 183: 672-679; Frakar et al. (1978) Biochem. Biophys. Res. Comm. 80:849-857; Chio et al. (1990) Nature 343: 266-269; Dahlman et al. (1988) Biochem 27: 1813-1817; Strader et al. (1989) J. Biol. Chem. 264: 13572-13578; and D'Dowd et al. (1988) J. Biol. Chem. 263: 15985-15992.

In a typical ligand/receptor binding assay useful in the practice of this invention, purified OP-1 having a known, quantifiable affinity for a pre-selected OP-1 receptor (see, for example, Ten Dijke et al. (1994) Science 264:101-103, the disclosure of which is incorporated herein by reference) is labeled with a detectable moiety, for example, a radiolabel, a chromogenic label, or a fluorogenic label. Aliquots of purified receptor, receptor binding domain fragments, or cells expressing the receptor of interest on their surface are incubated with labeled OP-1 in the presence of various concentrations of the unlabeled morphogen analog. The relative binding affinity of the morphogen analog may be measured by quantitating the ability of the candidate (unlabeled morphogen analog) to inhibit the binding of labeled OP-1 with the receptor. In performing the assay, fixed concentrations of the receptor and the OP-1 are incubated in the presence and absence of unlabeled morphogen analog. Sensitivity may be increased by preincubating the receptor with the candidate morphogen analog before adding labeled OP-1. After the labeled competitor has been added, sufficient time is allowed for adequate competitor binding, and then free and bound labeled OP-1 are separated from one another, and one or the other measured.

Labels useful in the practice of the screening procedures include radioactive labels (e.g., 125I, 131I, 111In or 77Br), chromogenic labels, spectroscopic labels (such as those disclosed in Haughland (1994) "Handbook of Fluorescent and Research Chemicals 5 ed." by Molecular Probes, Inc., Eugene, OR), or conjugated enzymes having high turnover rates, for example, horseradish peroxidase, alkaline phosphatase, or β-galactosidase, used in combination with chemiluminescent or fluorogenic substrates.

The biological activity, namely the agonist or antagonist properties of the resulting morphogen analogs subsequently may be characterized using any conventional *in vivo* and *in vitro* assays that have been developed to measure the biological activity of OP-1. A variety of specific assays believed to be useful in the practice of the invention are set forth in detail in Example 1, hereinbelow.

Furthermore, it is appreciated that many of the standard OP-1 assays may be automated thereby facilitating the screening of a large number of morphogen analogs at the same time. Such automation procedures are within the level of skill in the art of drug screening and, therefore, are not discussed herein.

Following the identification of useful morphogen analogs, the morphogenic analogs may be produced in commercially useful quantities (e.g., without limitation, gram and kilogram quantities), for example, by producing cell lines that express the morphogen analogs of interest or by producing synthetic peptides defining the appropriate amino acid sequence. It is appreciated, however, that conventional methodologies for producing the appropriate cell lines and for producing synthetic peptides are well known and thoroughly documented in the art, and so are not discussed in detail herein.

### VIII. Formulation and Bioactivity.

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Morphogen analogs, including OP-1 analogs, can be formulated for administration to a

25 mammal, preferably a human in need thereof as part of a pharmaceutical composition. The
composition can be administered by any suitable means, e.g., parenterally, orally or locally.

Where the morphogen analog is to be administered locally, as by injection, to a desired tissue
site, or systemically, such as by intravenous, subcutaneous, intramuscular, intraorbital,
ophthalmic, intraventricular, intracranial, intracapsular, intraspinal, intracisternal, intraperitoneal,

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buccal, rectal, vaginal, intranasal or aerosol administration, the composition preferably comprises an aqueous solution. The solution preferably is physiologically acceptable, such that administration thereof to a mammal does not adversely affect the mammal's normal electrolyte and fluid volume balance. The aqueous solution thus can comprise, e.g., normal physiologic saline (0.9% NaCl, 0.15M), pH 7-7.4.

Useful solutions for oral or parenteral systemic administration can be prepared by any of the methods well known in the pharmaceutical arts, described, for example, in "Remington's Pharmaceutical Sciences" (Gennaro, A., ed., Mack Pub., 1990, the disclosure of which is incorporated herein by reference). Formulations can include, for example, polyalkylene glycols such as polyethylene glycol, oils of vegetable origin, hydrogenated naphthalenes, and the like. Formulations for direct administration, in particular, can include glycerol and other compositions of high viscosity. Biocompatible, preferably bioresorbable polymers, including, for example, hyaluronic acid, collagen, tricalcium phosphate, polybutyrate, polylactide, polyglycolide and lactide/glycolide copolymers, may be useful excipients to control the release of the morphogen analog in vivo.

Other potentially useful parenteral delivery systems for the present analogs can include ethylene-vinyl acetate copolymer particles, osmotic pumps, implantable infusion systems, and liposomes. Formulations for inhalation administration can contain as excipients, for example, lactose, or can be aqueous solutions containing, for example, polyoxyethylene-9-lauryl ether, glycocholate or deoxycholate, or oily solutions for administration in the form of nasal drops or as a gel to be applied intranasally.

Alternatively, the morphogen analogs, including OP-1 analogs, identified as described herein may be administered orally. For example, liquid formulations of morphogen analogs can be prepared according to standard practices such as those described in "Remington's Pharmaceutical Sciences" (supra). Such liquid formulations can then be added to a beverage or another food supplement for administration. Oral administration can also be achieved using aerosols of these liquid formulations. Alternatively, solid formulations prepared using artrecognized emulsifiers can be fabricated into tablets, capsules or lozenges suitable for oral administration.

Optionally, the analogs can be formulated in compositions comprising means for enhancing uptake of the analog by a desired tissue. For example, tetracycline and diphosphonates (bisphosphonates) are known to bind to bone mineral, particularly at zones of bone remodeling, when they are provided systemically in a mammal. Accordingly, such components can be used to enhance delivery of the present analogs to bone tissue. Alternatively, an antibody or portion thereof that binds specifically to an accessible substance specifically associated with the desired target tissue, such as a cell surface antigen, also can be used. If desired, such specific targeting molecules can be covalently bound to the present analog, e.g., by chemical crosslinking or by using standard genetic engineering techniques to create, for example, an acid labile bond such as an Asp-Pro linkage. Useful targeting molecules can be designed, for example, according to the teachings of U.S. 5,091,513.

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It is contemplated also that some of the morphogen analogs may exhibit the highest levels of activity *in vivo* when combined with carrier matrices i.e., insoluble polymer matrices. See for example, U.S. Patent No. 5,266,683 the disclosure of which is incorporated by reference herein. Currently preferred carrier matrices are xenogenic, allogenic or autogenic in nature. It is contemplated, however, that synthetic materials comprising polylactic acid, polyglycolic acid, polybutyric acid, derivatives and copolymers thereof may also be used to generate suitable carrier matrices. Preferred synthetic and naturally derived matrix materials, their preparation, methods for formulating them with the morphogen analogs of the invention, and methods of administration are well known in the art and so are not discussed in detailed herein. See for example, U.S. Patent No. 5,266,683.

Still further, the present analogs can be administered to the mammal in need thereof either alone or in combination with another substance known to have a beneficial effect on tissue morphogenesis. Examples of such substances (herein, cofactors) include substances that promote tissue repair and regeneration and/or inhibit inflammation. Examples of useful cofactors for stimulating bone tissue growth in osteoporotic individuals, for example, include but are not limited to, vitamin D<sub>3</sub>, calcitonin, prostaglandins, parathyroid hormone, dexamethasone, estrogen and IGF-I or IGF-II. Useful cofactors for nerve tissue repair and regeneration can include nerve growth factors. Other useful cofactors include symptom-alleviating cofactors, including antiseptics, antibiotics, antiviral and antifungal agents, analgesics and anesthetics.

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Analogs preferably are formulated into pharmaceutical compositions by admixture with pharmaceutically acceptable, nontoxic excipients and carriers. As noted above, such compositions can be prepared for systemic, e.g., parenteral, administration, particularly in the form of liquid solutions or suspensions; for oral administration, particularly in the form of tablets or capsules; or intranasally, particularly in the form of powders, nasal drops or aerosols. Where adhesion to a tissue surface is desired, the composition can comprise a fibrinogen-thrombin dispersant or other bioadhesive such as is disclosed, for example, in PCT US91/09275, the disclosure of which is incorporated herein by reference. The composition then can be painted, sprayed or otherwise applied to the desired tissue surface.

The compositions can be formulated for parenteral or oral administration to humans or other mammals in therapeutically effective amounts, e.g., amounts which provide appropriate concentrations of the morphogen analog to target tissue for a time sufficient to induce the desired effect. Preferably, the present compositions alleviate or mitigate the mammal's need for a morphogen-associated biological response, such as maintenance of tissue-specific function or restoration of tissue-specific phenotype to senescent tissues (e.g., osteopenic bone tissue).

As will be appreciated by those skilled in the art, the concentration of the compounds described in a therapeutic composition will vary depending upon a number of factors, including the dosage of the drug to be administered, the chemical characteristics (e.g., hydrophobicity) of the compounds employed, and the route of administration. The preferred dosage of drug to be administered also is likely to depend on such variables as the type and extent of a disease, tissue loss or defect, the overall health status of the particular patient, the relative biological efficacy of the compound selected, the formulation of the compound, the presence and types of excipients in the formulation, and the route of administration. In general terms, the therapeutic molecules of this invention may be provided to an individual where typical doses range from about 10 ng/kg to about 1 g/kg of body weight per day; with a preferred dose range being from about 0.1 mg/kg to 100 mg/kg of body weight.

#### IX. Examples

Practice of the invention will be more fully understood from the following examples, which are presented herein for illustrative purposes only, and should not be construed as limiting the invention in any way.

# 5 Example 1. Introduction of Inter-chain Disulfide Bonds to Stabilize the hOP-1 Dimer.

As discussed in section V.A.(i) it is contemplated that introduction of one or more additional inter-chain disulfide may stabilize further the hOP-1 dimer. The introduction of additional inter-chain disulfide bonds is described here.

A Sma I to Bam HI fragment of the human OP-1 cDNA as described in Ozkaynak et al. (1990) supra is cloned into Bluescript KS+ (available from Stratagene Cloning Systems, La Jolla, CA), previously cleaved with Eco RV and Bam HI. Upon transformation into E. coli, the resulting colonies are screened by a blue-white selection process wherein the desired colonies containing the OP-1 cDNA insert are blue. The correct clone may be identified by restriction screening to give the following expected restriction fragments.

Restriction Enzyme	Fragment size (bp)
EcoR I	84, 789, 3425
Xho I	161, 1223, 2914
Sac II	97, 650, 3551

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In order to introduce two additional inter-chain disulfide bridges, a double cysteine mutant containing Asn 83 to Cys and Asn 130 to Cys replacements is produced. The cysteine mutant can be prepared by site-directed mutagenesis using synthetic oligonucleotides and either PCR or the site-directed mutagenesis methods, see for example, Kunkel et al. (1985) Proc. Natl. Acad. Sci. USA 822: 488; Kunkel et al. (1985) Meth. Enzymol. 154: 367 and U.S. Patent No. 4,873,192. Neither mutation causes a frameshift and, therefore, E. coli transformed with mutagenesis products that give white colonies indicate an error in the sequence. The presence of the appropriate mutation is verified by conventional dideoxy sequencing.

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Then, linkers are introduced into the N- and C-termini of the mutant gene by oligonucleotide-directed mutagenesis using appropriate oligonucleotides. A preferred N terminal linker introduces a unique Not I site and a preferred C terminal linker introduces a non-suppressible stop codon TAA at the end of the mutein gene followed by a unique Bgl II site (AGATCT). Each of the resulting mutant genes are excised from the cloning vector by the restriction enzymes Nde I and Bgl II, isolated, and ligated independently into pET vector (New England Biolabs, Beverly, MA) previously cleaved with Nde I and Bam HI. The ligation products then are transformed into *E. coli* and transformants containing, and expressing each individual mutant protein are identified.

Expression of the double cysteine containing mutant analog is induced after the expression of T7 RNA polymerase (initiated through infected with λCE6 phage). During expression, the mutant analog is produced as inclusion granules which are harvested from the cell paste. Then, the mutant protein is dissolved in 6M guanidine-HCl, 0.2M Tris-HCl, pH 8.2 and 0.1 M 2-mercaptoethanol, and the mixture dialyzed exhaustively against 6M urea, 2.5 mM Tris-HCl, pH 7.5 and 1 mM EDTA. 2-mercaptoethanol is added to a final concentration of 0.1M and the solution incubated at room temperature. The mixture is dialyzed exhaustively against buffer containing 2.5 mM Tris-HCl, pH 7.5 and 1mM EDTA. Folded mutant protein is purified by affinity chromatography on a column packed with surface immobilized OP-1 receptor. Unbound material is removed by washing as described above and the specific OP-1 receptor binding material eluted.

Following purification the stabilizing effect of the additional bond is determined by fluorescence polarization. For example, the rotational rates of morphogen analog (mutein) and natural hOP-1 are determined as a function of temperature using a fluorescence spectrophotometer modified for fluorescence anisotropy (Photon Technology International). It is anticipated that the mutein dimer will exhibit a lower rational rate upto a higher temperature than natural hOP-1 dimer, thereby indicating that the mutein dimer remains as a dimer and is more stable upto a higher temperature than is the wild type protein.

The biological activity of the resulting mutant protein or mutein can be tested using any of the bioassays developed to date for determining the biological activity of native hOP-1. A variety of such exemplary assays are described below. The assays which follow are recited for

ease of testing. Specific in vivo assays for testing the efficacy of a morphogenic protein or analog in an application to repair or regenerate damaged bone, liver, kidney, or nerve tissue, periodontal tissue, including cementum and/or periodontal ligament, gastrointestinal and renal tissues, and immune-cell mediated damages tissues are disclosed in publicly available documents, which include, for example, EP 0575,555; WO93/04692; WO93/05751; WO/06399; WO94/03200; WO94/06449; and WO94/06420. The skilled artisan can test an analog in any of these assays without undue experimentation.

#### A. Mitogenic Effect on Rat and Human Osteoblasts

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The following example is a typical assay useful in determining whether an OP-1 morphogen analog induces proliferation of osteoblasts in vitro. It is contemplated that in this, and all other examples using osteoblast cultures, preferably uses rat osteoblast-enriched primary cultures. Although these cultures are heterogeneous in that the individual cells are at different stages of differentiation, the culture is believed to more accurately reflect the metabolism and function of osteoblasts in vivo than osteoblast cultures obtained from established cell lines. Unless otherwise indicated, all chemicals referenced are standard, commercially available reagents, readily available from a number of sources, including Sigma Chemical, Co., St. Louis; Calbiochem, Corp., San Diego and Aldrich Chemical Co., Milwaukee.

Briefly, rat osteoblast-enriched primary cultures are prepared by sequential collagenase digestion of newborn rat calvaria (e.g., from 1-2 day-old animals, Long-Evans strain, Charles River Laboratories, Wilmington, MA), following standard procedures, such as are described, for example, in Wong et al. (1975) Proc. Natl. Acad. Sci. USA 72: 3167-3171. Rat osteoblast single cell suspensions then are plated onto a multi-well plate (e.g., a 24 well plate at a concentration of 50,000 osteoblasts per well) in alpha MEM (modified Eagle's medium, Gibco, Inc., NY) containing 10% FBS (fetal bovine serum), L-glutamine and penicillin/streptomycin. The cells are incubated for 24 hours at 37°C, at which time the growth medium is replaced with alpha MEM containing 1% FBS and the cells incubated for an additional 24 hours so that cells are in serum-deprived growth medium at the time of the experiment.

The cultured cells are divided into four groups: (1) wells which receive, for example, 0.1, 1.0, 10.0, 40.0 and 80.0 ng of the OP-1 morphogen analog (mutein), (2) wells which receive 0.1,

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1.0, 10.0 and 40.0 ng of wild type OP-1; (3) wells which receives 0.1, 1.0, 10.0, and 40.0 ng of TGF-β, and (4) the control group, which receive no growth factors. The cells then are incubated for an additional 18 hours after which the wells are pulsed with 2mCi/well of <sup>3</sup>H-thymidine and incubated for six more hours. The excess label then is washed off with a cold solution of 0.15 M NaCl and then 250 ml of 10% tricholoracetic acid is added to each well and the wells incubated at room temperature for 30 minutes. The cells then are washed three times with cold distilled water, and lysed by the addition of 250 ml of 1% sodium dodecyl sulfate (SDS) for a period of 30 minutes at 37°C. The resulting cell lysates are harvested using standard means and the incorporation of <sup>3</sup>H-thymidine into cellular DNA determined by liquid scintillation as an indication of mitogenic activity of the cells. In the experiment, it is contemplated that the OP-1 morphogen analog construct (mutein), like natural OP-1, will stimulate <sup>3</sup>H-thymidine incorporation into DNA, and therefore promote osteoblast cell proliferation. In contrast, the effect of the TGF-β is expected to be transient and biphasic. Furthermore, it is contemplated that at higher concentrations, TGF-β will have no significant effect on osteoblast cell proliferation.

The *in vitro* effect of the OP-1 morphogen analog on osteoblast proliferation also may be evaluated using human primary osteoblasts (obtained from bone tissue of a normal adult patient and prepared as described above) and on human osteosarcoma-derived cell lines.

#### B. Progenitor Cell Stimulation.

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The following example is designed to demonstrate the ability of OP-1 morphogen analogs to stimulate the proliferation of mesenchymal progenitor cells. Useful naive stem cells include pluripotent stem cells, which may be isolated from bone marrow or umbilical cord blood using conventional methodologies, (see, for example, Faradji et al. (1988) Vox Sang. 55 (3): 133-138 or Broxmeyer et al. (1989) Proc. Natl. Acad. Sci. USA. 86: 3828-3832), as well as naive stem cells obtained from blood. Alternatively, embryonic cells (e.g., from a cultured mesodermal cell line) may be used.

Another method for obtaining progenitor cells and for determining the ability of OP-1 morphogen analogs to stimulate cell proliferation is to capture progenitor cells from an *in vivo* source. For example, a biocompatible matrix material able to allow the influx of migratory progenitor cells may be implanted at an *in vivo* site long enough to allow the influx of migratory

progenitor cells. For example, a bone-derived, guanidine-extracted matrix, formulated as disclosed for example in Sampath *et al.* (1983) *Proc. Natl. Acad. Sci. USA 80*: 6591-6595, or U.S. Patent No. 4,975,526, may be implanted into a rat at a subcutaneous site, essentially following the method of Sampath *et al.* After three days the implant is removed, and the progenitor cells associated with the matrix dispersed and cultured.

Progenitor cells, however obtained, then are incubated *in vitro* with the candidate OP-1 morphogen analog under standard cell culture conditions, such as those described hereinbelow. In the absence of external stimuli, the progenitor cells do not, or only minimally, proliferate on their own in culture.

10 However, progenitor cells cultured in the presence of a biologically active OP-1 morphogen analog, like OP-1, will proliferate. Cell growth can be determined visually or spectrophotometrically using standard methods well known in the art.

#### C. Morphogen-Induced Cell Differentiation.

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A variety of assays also can be used to determine OP-1 based morphogen analog-induced cellular differentiation.

#### (1) Embryonic Mesenchyme Differentiation

As with natural OP-1, it is contemplated that the OP-1 morphogen analog (mutein) can induce cell differentiation. The ability of OP-1 morphogen analogs to induce cell differentiation can be demonstrated by culturing early mesenchymal cells in the presence of OP-1 morphogen analog and then studying the histology of the cultured cells by staining with toluidine blue using standard cell culturing and cell staining methodologies well described in the art. For example, it is known that rat mesenchymal cells destined to become mandibular bone, when separated from the overlying epithelial cells at stage 11 and cultured *in vitro* under standard tissue culture conditions, e.g., in a chemically defined, serum-free medium, containing for example, 67% DMEM (Dulbecco's modified Eagle's medium), 22% F-12 medium, 10mM Hepes pH 7, 2mM glutamine, 50 mg/ml transferrin, 25 mg/ml insulin, trace elements, 2mg/ml bovine serum albumin coupled to oleic acid, with HAT (0.1 mM hypoxanthine, 10mM aminopterin, 12 mM thymidine, will not continue to differentiate. However, if these same cells are left in contact with the overlying endoderm for an additional day, at which time they become stage 12 cells, they

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will continue to differentiate on their own *in vitro* to form chondrocytes. Further differentiation into osteoblasts and, ultimately, mandibular bone, requires an appropriate local environment, e.g., a vascularized environment.

It is anticipated that, as with natural OP-1, stage 11 mesenchymal cells, cultured *in vitro* in the presence of OP-1 morphogen analog (mutein), e.g., 10-100 ng/ml, will continue to differentiate *in vitro* to form chondrocytes just as they continue to differentiate *in vitro* if they are cultured with the cell products harvested from the overlying endodermal cells. This experiment can be performed with different mesenchymal cells to demonstrate the cell differentiation capability of OP-1 morphogen analog in different tissues.

As another example of morphogen-induced cell differentiation, the ability of OP-1 morphogen analogs to induce osteoblast differentiation can be demonstrated *in vitro* using primary osteoblast cultures, or osteoblast-like cells lines, and assaying for a variety of bone cell markers that are specific markers for the differentiated osteoblast phenotype, e.g., alkaline phosphatase activity, parathyroid hormone-mediated cyclic AMP (cAMP) production, osteocalcin synthesis, and enhanced mineralization rates.

# (2) Induction of Alkaline Phosphatase Activity in Osteoblasts.

Cultured osteoblasts in serum-free medium are incubated with a range of OP-1 morphogen analog concentrations, for example, 0.1, 1.0, 10.0, 40.0 or 80.0 ng OP-1 morphogen analog/ml medium; or with a similar concentration range of natural OP-1 or TGF-β. After a 72 hour incubation the cell layer is extracted with 0.5 ml of 1% Triton X-100. The resultant cell extract is centrifuged, and 100 ml of the extract is added to 90 ml of para-nitrosophenylphosphate (PNPP)/glycerine mixture and incubated for 30 minutes in a 37°C water bath and the reaction stopped with 100 ml NaOH. The samples then are run through a plate reader (e.g., Dynatech MR700 plate reader, and absorbance measured at 400 nm, using p-nitrophenol as a standard) to determine the presence and amount of alkaline phosphate activity. Protein concentrations are determined by the BioRad method. Alkaline phosphatase activity is calculated in units/mg protein, where 1 unit=1 nmol p-nitrophenol liberated/30 minutes at 37°C.

It is contemplated that the OP-1 morphogen analog, like natural OP-1, will stimulate the production of alkaline phosphatase in osteoblasts thereby promoting the growth and expression

of the osteoblast differentiated phenotype. The long term effect of OP-1 morphogen analog on the production of alkaline phosphatase by rat osteoblasts also can be demonstrated as follows.

Rat osteoblasts are prepared and cultured in multi-well plates as described above. In this example six sets of 24 well plates are plated with 50,000 rat osteoblasts per well. The wells in each plate, prepared as described above, then are divided into three groups: (1) those which receive, for example, 1 ng of OP-1 morphogen analog per ml of medium; (2) those which receive 40 ng of OP-1 morphogen analog per ml of medium; and (3) those which receive 80 ng of OP-1 morphogen analog per ml of medium. Each plate then is incubated for different lengths of time: 0 hours (control time), 24 hours, 48 hours, 96 hours, 120 hours and 144 hours. After each incubation period, the cell layer is extracted with 0.5 ml of 1% Triton X-100. The resultant cell extract is centrifuged, and alkaline phosphatase activity determined using para-nitrosophenylphosphate (PNPP), as above. It is contemplated that the OP-1 morphogen analog, like natural OP-1, will stimulate the production of alkaline phosphatase in osteoblasts in a dosedependent manner so that increasing doses of OP-1 morphogen analog will further increase the level of alkaline phosphatase production. Moreover, it is contemplated that the OP-1 morphogen analog-stimulated elevated levels of alkaline phosphatase in the treated osteoblasts will last for an extended period of time.

#### (3) Induction of PTH-Mediated cAMP.

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This experiment is designed to test the effect of OP-1 morphogen analogs on parathyroid hormone-mediated cAMP production in rat osteoblasts *in vitro*. Briefly, rat osteoblasts are prepared and cultured in a multiwell plate as described above. The cultured cells then are divided into four groups: (1) wells which receive, for example, 1.0, 10.0 and 40.0 ng OP-1 morphogen analog/ml medium); (2) wells which receive for example, natural OP-1, at similar concentration ranges; (3) wells which receive for example, TGF-β, at similar concentration ranges; and (4) a control group which receives no growth factors. The plate then is incubated for another 72 hours. At the end of the 72 hours the cells are treated with medium containing 0.5% bovine serum albumin (BSA) and 1mM 3-isobutyl-1-methylxanthine for 20 minutes followed by the addition into half of the wells of human recombinant parathyroid hormone (hPTH, Sigma, St. Louis) at a concentration of 200 ng/ml for 10 minutes. The cell layer then is extracted from each well with 0.5 ml of 1% Triton X-100. The cAMP levels then are determined using a

radioimmunoassay kit (e.g., Amersham, Arlington Heights, Illinois). It is contemplated that OP-1 morphogen analog alone, like OP-1, will stimulate an increase in the PTH-mediated cAMP response, thereby promoting the growth and expression of the osteoblast differentiated phenotype.

# (4) Induction of Osteocalcin Production.

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Osteocalcin is a bone-specific protein synthesized by osteoblasts which plays an integral role in the rate of bone mineralization *in vivo*. Circulating levels of osteocalcin in serum are used as a marker for osteoblast activity and bone formation *in vivo*. Induction of osteocalcin synthesis in osteoblast-enriched cultures can be used to demonstrate OP-1 morphogen analog efficacy *in vitro*.

Rat osteoblasts are prepared and cultured in a multi-well plate as above. In this experiment the medium is supplemented with 10%FBS, and on day 2, cells are fed with fresh medium supplemented with fresh 10 mM  $\beta$ -glycerophosphate (Sigma, Inc.). Beginning on day 5 and twice weekly thereafter, cells are fed with a complete mineralization medium containing all of the above components plus fresh L(+)-ascorbate, at a final concentration of 50mg/ml medium. OP-1 morphogen analog then is added to the wells directly, e.g., in 50% acetonitrile (or 50% ethanol) containing 0.1% trifluoroacetic acid (TFA), at no more than 5ml morphogen analog/ml medium. Control wells receive solvent vehicle only. The cells then are re-fed and the conditioned medium sample diluted 1:1 in standard radioimmunoassay buffer containing standard protease inhibitors and stored at -20°C until assayed for osteocalcin. Osteocalcin synthesis is measured by standard radioimmunoassay using a commercially available osteocalcin-specific antibody.

Mineralization is determined on long term cultures (13 day) using a modified von Kossa staining technique on fixed cell layers: cells are fixed in fresh 4% paraformaldehyde at 23°C for 10 min, following rinsing cold 0.9% NaCl. Fixed cells then are stained for endogenous alkaline phosphatase at pH 9.5 for 10 min, using a commercially available kit (Sigma, Inc.). Purple stained cells then are dehydrated with methanol and air dried. After 30 min incubation in 3% AgNO3 in the dark, H2O-rinsed samples are exposed for 30 sec to 254 nm UV light to develop

the black silver-stained phosphate nodules. Individual mineralized foci (at least 20 mm in size) are counted under a dissecting microscope and expressed as nodules/culture.

It is contemplated that the OP-1 morphogen analog, like natural OP-1, will stimulate osteocalcin synthesis in osteoblast cultures. Furthermore, it is contemplated that the increased osteocalcin synthesis in response to OP-1 morphogen analog will be in a dose dependent manner thereby showing a significant increase over the basal level after 13 days of incubation. Enhanced osteocalcin synthesis also can be confirmed by detecting the elevated osteocalcin mRNA message (20-fold increase) using a rat osteocalcin-specific probe. In addition, the increase in osteocalcin synthesis correlates with increased mineralization in long term osteoblast cultures as determined by the appearance of mineral nodules. It is contemplated also that OP-1 morphogen analog, like natural OP-1, will increase significantly the initial mineralization rate as compared to untreated cultures.

### (5) Morphogen-Induced CAM Expression

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Members of the BMP/OP family (see Figure 6) induce CAM expression, particularly N-CAM expression, as part of their induction of morphogenesis (see copending U.S.S.N. 922,813). CAMs are morphoregulatory molecules identified in all tissues as an essential step in tissue development. N-CAMs, which comprise at least 3 isoforms (N-CAM-180, N-CAM-140 and N-CAM-120, where "180", "140" and "120" indicate the apparent molecular weights of the isoforms as measured by SDS polyacrylamide gel electrophoresis) are expressed at least transiently in developing tissues, and permanently in nerve tissue. Both the N-CAM-180 and N-CAM-140 isoforms are expressed in both developing and adult tissue. The N-CAM-120 isoform is found only in adult tissue. Another neural CAM is L1.

The ability of OP-1 based morphogen analogs to stimulate CAM expression may be demonstrated using the following protocol, using NG108-15 cells. NG108-15 is a transformed hybrid cell line (neuroblastoma x glioma, America Type Culture Collection (ATCC), Rockville, MD), exhibiting a morphology characteristic of transformed embryonic neurons. As described in Example D, below, untreated NG108-15 cells exhibit a fibroblastic, or minimally differentiated, morphology and express only the 180 and 140 isoforms of N-CAM normally associated with a developing cell. Following treatment with members of the vg/dpp subgroup these cells exhibit a

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morphology characteristic of adult neurons and express enhanced levels of all three N-CAM isoforms.

In this example, NG108-15 cells are cultured for 4 days in the presence of increasing concentrations of either the OP-1 morphogen analog or natural OP-1 using standard culturing procedures, and standard Western blots are performed on whole cell extracts. N-CAM isoforms are detected with an antibody which crossreacts with all three isoforms, mAb H28.123, obtained from Sigma Chemical Co., St. Louis, the different isoforms being distinguishable by their different mobilities on an electrophoresis gel. Control NG108-15 cells (untreated) express both the 140 kDa and the 180 kDa isoforms, but not the 120 kDa, as determined by Western blot analyses using up to 100 mg of protein. It is contemplated that treatment of NG108-15 cells with OP-1 morphogen analog, like natural OP-1 may result in a dose-dependent increase in the expression of the 180 kDa and 140 kDa isoforms, as well as the induction of the 120 kDa isoform. In addition, it is contemplated that the OP-1 morphogen analog, like natural OP-1-induced CAM expression may correlate with cell aggregation, as determined by histology.

# (D) OP-1 Morphogen Analog-Induced Redifferentiation of Transformed Phenotype

It is contemplated that OP-1 morphogen analog, like natural OP-1, also induces redifferentiation of transformed cells to a morphology characteristic of untransformed cells. The examples provided below detail morphogen-induced redifferentiation of a transformed human cell line of neuronal origin (NG108-15); as well as mouse neuroblastoma cells (N1E-115), and human embryo carcinoma cells, to a morphology characteristic of untransformed cells.

As described above, NG108-15 is a transformed hybrid cell line produced by fusing neuroblastoma x glioma cells (obtained from ATCC, Rockville, MD), and exhibiting a morphology characteristic of transformed embryonic neurons, e.g., having a fibroblastic morphology. Specifically, the cells have polygonal cell bodies, short, spike-like processes and make few contacts with neighboring cells. Incubation of NG108-15 cells, cultured in a chemically defined, serum-free medium, with 0.1 to 300 ng/ml of morphology analog or natural OP-1 for four hours induces an orderly, dose-dependent change in cell morphology.

For example, NG108-15 cells are subcultured on poly-L-lysine coated 6 well plates. Each well contains 40-50,000 cells in 2.5 ml of chemically defined medium. On the third day,

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2.5 ml of OP-1 morphogen analog or natural OP-1 in 60% ethanol containing 0.025% trifluoroacetic is added to each well. The media is changed daily with new aliquots of morphogen. It is contemplated that OP-1 morphogen analog, like OP-1, may induce a dose-dependent redifferentiation of the transformed cells, including a rounding of the soma, an increase in phase brightness, extension of the short neurite processes, and other significant changes in the cellular ultrastructure. After several days it is contemplated also that treated cells may begin to form epithelioid sheets that then become highly packed, multi-layered aggregates, as determined visually by microscopic examination.

Moreover, it is contemplated that the redifferentiation may occur without any associated changes in DNA synthesis, cell division, or cell viability, making it unlikely that the morphologic changes are secondary to cell differentiation or a toxic effect of the morphogen. In addition, it is contemplated that the morphogen analog-induced redifferentiation may not inhibit cell division, as determined by <sup>3</sup>H-thymidine uptake, unlike other molecules such as butyrate, DMSO, retinoic acid or Forskolin, which have been shown to stimulate differentiation of transformed cells in analogous experiments. Thus, it is contemplated that the OP-1 morphogen analog, like natural OP-1, may maintain cell stability and viability after inducing redifferentiation.

The morphogen described herein would, therefore, provide useful therapeutic agents for the treatment of neoplasias and neoplastic lesions of the nervous system, particularly in the treatment of neuroblastomas, including retinoblastomas, and gliomas.

#### (E) Maintenance of Phenotype.

OP-1 morphogen analogs, like natural OP-1, also may be used to maintain a cell's differentiated phenotype. This application is particularly useful for inducing the continued expression of phenotype in senescent or quiescent cells.

## (1) In Vitro Model for Phenotypic Maintenance

The phenotypic maintenance capability of morphogens is determined readily. A number of differentiated cells become senescent or quiescent after multiple passages in vitro under standard tissue culture conditions well described in the art (e.g., Culture of Animal Cells: A Manual of Basic Techniques, C.R. Freshney, ed., Wiley, 1987). However, if these cells are

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cultivated *in vitro* in association with a morphogen such as OP-1, cells are stimulated to maintain expression of their phenotype through multiple passages. For example, the alkaline phosphatase activity of cultured osteoblasts, such as cultured osteosarcoma cells and calvaria cells, is significantly reduced after multiple passages *in vitro*. However, if the cells are cultivated in the presence of OP-1, alkaline phosphatase activity is maintained over extended periods of time. Similarly, phenotypic expression of myocytes also is maintained in the presence of a morphogen. In the experiment, osteoblasts are cultured as described in Example A. The cells are divided into groups, incubated with varying concentrations of either OP-1 morphogen analog or natural OP-1 (e.g., 0-300 ng/ml) and passaged multiple times (e.g., 3-5 times) using standard methodology. Passaged cells then are tested for alkaline phosphatase activity, as described in Example C as an indication of differentiated cell metabolic function. It is contemplated that osteoblasts cultured in the absence of OP-1 morphogen analog may have reduced alkaline phosphatase activity, as compared to OP-1 morphogen analog, or natural OP-1-treated cells.

# (2) In Vivo Model for Phenotypic Maintenance.

Phenotypic maintenance capability also may be demonstrated *in vivo*, using a standard rat model for osteoporosis. Long Evans female rats (Charles River Laboratories, Wilmington, MA) are sham-operated (control animals) or ovariectomized using standard surgical techniques to produce an osteoporotic condition resulting from decreased estrogen production. Following surgery, e.g., 200 days after ovariectomy, rats are systemically provided with phosphate buffered saline (PBS) or morphogen, (e.g., OP-1 morphogen analog, or natural OP-1, 1-100 mg) for 21 days (e.g., by daily tail vein injection.) The rats then are sacrificed and serum alkaline phosphatase levels, serum calcium levels, and serum osteocalcin levels are determined, using standard methodologies as described therein and above. It is contemplated that the OP-1 morphogen analog treated rats, like the OP-1 treated rats may exhibit elevated levels of osteocalcin and alkaline phosphatase activity. It is contemplated also that histomorphometric analysis on the tibial diaphyseal bone may show improved bone mass in OP-1 morphogen analog-treated animals as compared with untreated, ovariectomized rats.

### F. Proliferation of Progenitor Cell Populations

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Progenitor cells may be stimulated to proliferate in vivo or ex vivo. It is contemplated that cells may be stimulated in vivo by injecting or otherwise providing a sterile preparation containing the OP-1 morphogen analog into the individual. For example, the hematopoietic pluripotential stem cell population of an individual may be stimulated to proliferate by injecting or otherwise providing an appropriate concentration of OP-1 morphogen analog to the individual's bone marrow.

Progenitor cells may be stimulated *ex vivo* by contacting progenitor cells of the population to be enhanced with a morphogenically active OP-1 morphogen analog under sterile conditions at a concentration and for a time sufficient to stimulate proliferation of the cells. Suitable concentrations and stimulation times may be determined empirically, essentially following the procedure described in Example A, above. It is contemplated that a OP-1 morphogen analog concentration of between about 0.1-100 ng/ml and a stimulation period of from about 10 minutes to about 72 hours, or, more generally, about 24 hours, typically should be sufficient to stimulate a cell population of about 10<sup>4</sup> to 10<sup>6</sup> cells. The stimulated cells then may be provided to the individual as, for example, by injecting the cells to an appropriate *in vivo* locus. Suitable biocompatible progenitor cells may be obtained by any of the methods known in the art or described hereinabove.

### G. Regeneration of Damaged or Diseased Tissue

It is contemplated that OP-1 morphogen analogs may be used to repair diseased or damaged mammalian tissue. The tissue to be repaired preferably is assessed first, and excess necrotic or interfering scar tissue removed as needed, e.g., by ablation or by surgical, chemical, or other methods known in the medical arts.

OP-1 morphogen analog then may be provided directly to the tissue locus as part of a sterile, biocompatible composition, either by surgical implantation or injection. The morphogen analog also may be provided systemically, as by oral or parenteral administration. Alternatively, a sterile, biocompatible composition containing progenitor cells stimulated by a morphogenically active OP-1 morphogen analog may be provided to the tissue locus. The existing tissue at the

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locus, whether diseased or damaged, provides the appropriate matrix to allow the proliferation and tissue-specific differentiation of progenitor cells. In addition, a damaged or diseased tissue locus, particularly one that has been further assaulted by surgical means, provides a morphogenically permissive environment. Systemic provision of OP-1 morphogen analog may be sufficient for certain applications (e.g., in the treatment of osteoporosis and other disorders of the bone remodeling cycle).

In some circumstances, particularly where tissue damage is extensive, the tissue may not be capable of providing a sufficient matrix for cell influx and proliferation. In these instances, it may be necessary to provide progenitor cells stimulated by the OP-1 morphogen analog to the tissue locus in association with a suitable, biocompatible, formulated matrix, prepared by any of the means described below. The matrix preferably is *in vivo* biodegradable. The matrix also may be tissue-specific and/or may comprise porous particles having dimensions within the range of 70-850 µm, most preferably 150-420 µm.

OP-1 morphogen analog also may be used to prevent or substantially inhibit immune/inflammatory response-mediated tissue damage and scar tissue formation following an injury. OP-1 morphogen analog may be provided to a newly injured tissue locus, to induce tissue morphogenesis at the locus, preventing the aggregation of migrating fibroblasts into non-differentiated connective tissue. Preferably the OP-1 morphogen analog may be provided as a sterile pharmaceutical preparation injected into the tissue locus within five hours of the injury. Where an immune/inflammatory response is unavoidably or deliberately induced, as part of, for example, a surgical or other aggressive clinical therapy, OP-1 morphogen analog preferably may be provided prophylactically to the patient prior to, or concomitant with, the therapy.

Described below is a protocol for demonstrating whether a OP-1 morphogen analoginduces tissue morphogenesis in bone.

# (1) OP-1 Morphogen Analog-Induced Bone Morphogenesis.

A particularly useful mammalian tissue model system for demonstrating and evaluating the morphogenic activity of a morphogen analog is the endochondral bone tissue morphogenesis model known in the art and described, for example, in U.S. Pat. No. 4,968,590, incorporated herein by reference. The ability to induce endochondral bone formation includes the ability to

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induce proliferation and differentiation of progenitor cells into chondroblasts and osteoblasts, the ability to induce cartilage matrix formation, cartilage calcification, and bone remodeling, and the ability to induce formation of an appropriate vascular supply and hematopoietic bone marrow differentiation.

The local environment in which the morphogenic material is placed is important for tissue morphogenesis. As used herein, "local environment" is understood to include the tissue structural matrix and the environment surrounding the tissue. For example, in addition to needing an appropriate anchoring substratum for their proliferation, the cells stimulated by morphogens need signals to direct the tissue-specificity of their differentiation. These signals vary for the different tissues and may include cell surface markers. In addition, vascularization of new tissue requires a local environment which supports vascularization.

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The following sets forth various procedures for evaluating the *in vivo* morphogenic utility of OP-1 morphogen analogs and OP-1 morphogen analog containing compositions. The compositions may be injected or surgically implanted in a mammal, following any of a number of procedures well known in the art. For example, surgical implant bioassays may be performed essentially following the procedure of Sampath *et al.* (1983) *Proc. Natl. Acad. Sci. USA 80*: 6591-6595 and U.S. Pat No. 4,968,590.

Histological sectioning and staining is preferred to determine the extent of morphogenesis in vivo, particularly in tissue repair procedures. Excised implants are fixed in Bouins Solution, embedded in paraffin, and cut into 6-8 µm sections. Staining with toluidine blue or hemotoxylin/eosin demonstrates clearly the ultimate development of the new tissue. Twelve day implants are usually sufficient to determine whether the implants contain newly induced tissue.

Successful implants exhibit a controlled progression through the stages of induced tissue development allowing one to identify and follow the tissue-specific events that occur. For example, in endochondral bone formation the stages include: (1) leukocytes on day one; (2) mesenchymal cell migration and proliferation on days two and three; (3) chondrocyte appearance on days five and six; (4) cartilage matrix formation on day seven; (5) cartilage calcification on day eight; (6) vascular invasion, appearance of osteoblasts, and formation of new bone on days nine and ten; (7) appearance of osteoclastic cells, and the commencement of bone

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remodeling and dissolution of the implanted matrix on days twelve to eighteen; and (8) hematopoietic bone marrow differentiation in the resulting ossicles on day twenty-one.

In addition to histological evaluation, biological markers may be used as markers for tissue morphogenesis. Useful markers include tissue-specific enzymes whose activities may be assayed (e.g., spectrophotometrically) after homogenization of the implant. These assays may be useful for quantitation and for rapidly obtaining an estimate of tissue formation after the implants are removed from the animal. For example, alkaline phosphatase activity may be used as a marker for osteogenesis.

Incorporation of systemically provided OP-1 morphogen analog may be followed using labeled protein (e.g., radioactively labeled) and determining its localization in the new tissue, and/or by monitoring their disappearance from the circulatory system using a standard labeling protocol and pulse-chase procedure. OP-1 morphogen analog also may be provided with a tissue-specific molecular tag, whose uptake may be monitored and correlated with the concentration of OP-1 morphogen analog provided. As an example, ovary removal in female rats results in reduced bone alkaline phosphatase activity, and renders the rats predisposed to osteoporosis (as described in Example E). If the female rats now are provided with OP-1 morphogen analog, a reduction in the systemic concentration of calcium may be seen, which correlates with the presence of the provided OP-1 morphogen analog and which is anticipated to correspond with increased alkaline phosphatase activity.

# 20 Example 2. Enhancing the Solubility of a hOP-1 Dimer.

As described in section V.A.(ii), *supra*, it is contemplated that the solubility of the hOP-1 dimer can be enhanced by replacing hydrophobic amino acid residues located at the solvent accessible surface of hOP-1 dimer with more polar or hydrophilic amino acid residues. This example provides a description of such an approach.

A Sma I to Bam HI fragment of the human OP-1 cDNA as described in Ozkaynak et al. (1990) supra is cloned into a vector to produce a plasmid similar to the plasmid called pW24 in International Application PCT/US94/12063, the disclosure of which is incorporated herein by reference. The pW24 plasmid contains OP-1 cDNA under the transcriptional control of the CMV (cytomegalovirus) immediate early promoter. The selective marker on pW24 is the

neomycin gene which provides resistance to the cytostatic drug G418. The pW24 plasmid also employs an SV40 origin of replication (ori). The early SV40 promoter is used to drive transcription of the neomycin marker gene.

Then, the alanine at position 63 is mutated to a serine by site-directed mutagenesis using, for example, synthetic oligonucleotides and either PCR or the site-directed mutagenesis methods. See, for example, Kunkel et al. (1985) Proc. Natl. Acad. Sci. USA 822: 488; Kunkel et al. (1985) Meth. Enzymol. 154: 367 and U.S. Patent No. 4,873,192. The resulting mutation is confirmed by dideoxy sequencing.

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Two additional vectors have been developed for use in a triple transfection procedure along with pW24 to enhance OP-1 expression. One of the vectors employs the adenovirus E1A gene under the VA1 gene as translation stimulation for the gene DHFR gene. The other vector employs the adenovirus E1A gene under the control of the thymidine kinase promoter as a transactivating transcription activator. Both additional vectors, known as pH1130 and pH1176, as well as preferred transfection and screening procedures are described in International Application PCT/US94/12063.

Briefly, triple transfections are performed using the calcium phosphate coprecipitation procedure. CHO cells are cultured in  $\alpha$ MEM, containing 5% or 10% fetal bovine serum (FBS), non-essential amino acids, glutamine and antibiotics: penicillin and streptomycin. Stable cell line transfections are carried out by seeding 1-2x10<sup>6</sup> cells in a 9 cm. petri dish. Following an incubation period of up to 24-hour, each petri dish is transfected with 10-30 µg total vector DNA in equimolar amounts, by calcium phosphate coprecipitation followed by glycerol shock using standard methodology. Cells are incubated at 37°C in growth medium for 24 hours, then transferred to selection medium. All cultures are fed once or twice weekly with fresh selective medium. After 10 - 21 days, resistant colonies are picked and assayed for protein production.

Approximately 30 individual clones are selected, transferred to a 24-well petri dish, and allowed to grow to confluence in serum-containing media. The conditioned media from all surviving clones is screened for protein production using a standard ELISA (enzyme-linked immunosorbent assay) or Western blot. The methodologies for these assay protocols as well as

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for generating antibodies for use in these assays are well described in the art (see e.g., Ausubel, supra).

Under such conditions, the VA1 and E1A genes typically act synergistically to enhance OP1 expression in unamplified transfected CHO cells. Candidate cell lines identified by the screening protocol, then are seeded on ten 100 mm petri dishes at a cell density of either 50 or 100 cells per plate, and with a higher drug concentration (e.g., 1.0-µm).

After 10-21 days of growth, the clones are isolated using cloning cylinders and standard procedures, and cultured in 24-well plates. Then, clones are screened for OP-1 expression by Western immunoblots using standard procedures, and OP-1 expression levels compared to parental lines. Candidate cells showing higher protein production than cells of parental lines then are replated and grown in the presence of a still higher drug concentration (e.g., 5-20µm). Generally, no more than 2-3 rounds of these "amplification" cloning steps are necessary to achieve cell lines with high protein productivity. Useful high producing cell lines may be further subcloned to improve cell line homogeneity and product stability.

A currently preferred method of large scale protein production e.g., at least 2 liters, is by suspension culture of the host Chinese hamster ovary (CHO) cells. CHO cells prefer attachment but can be adapted to grow in suspension mode of cultivation. The cells are trypsinized from a culture dish, introduced to growth media containing 10% FBS and completely suspended to produce a single cell suspension. The single cell suspension is introduced to a spinner flask and placed in a 37°C 95% air/5% CO<sub>2</sub> humidified incubator. Over a period of time the cells are subcultured in medium with descending concentrations of serum.

Specifically, the adapted cells are introduced into a 3L spinner flask at an initial viable cell density of approximately  $2x10^5$  cells/ml. Preferred culture medium is DMEM/F-12 (1:1) (GIBCO, New York) supplemented with 2% FBS, and preferred agitation is approximately 50-60 rpm with a paddle impeller. After 7 days, the culture media is harvested, centrifuged at 1500 rpm and the clarified conditioned media stored at 4°C.

A representative purification scheme for purifying recombinant morphogenic protein involves three chromatographic steps (S-Sepharose, phenyl-Sepharose and C-18 HPLC) and is described in International Application PCT/US94/12063. Morphogen analog containing culture

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media is diluted to 6M urea, 0.05M NaC1, 13mM HEPES, pH 7.0 and loaded onto an S-Sepharose column, which acts as a strong cation exchanger. The column subsequently is developed with two salt elutions. The first elution employs a solution containing 0.1M NaC1, and the second elution employs a buffer containing 6M urea, 0.3M NaC1, 20mM HEPES, pH 7.0.

Ammonium sulfate is added to the 0.3M NaC1 fraction to give a solution containing 6M urea, 1M (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 0.3M NaC1, 20mM HEPES, pH 7.0. Then, the sample is loaded onto a phenyl-Sepharose column in the presence of 1M (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>). Then, the column is developed with two step elutions using decreasing concentrations of ammonium sulfate. The first elution employs 0.6M (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> and the second elution employs 6M urea, 0.3M NaC1, 20mM HEPES, pH 7.0 buffer. The material harvested from the second elution is dialyzed against water, followed by 30% acetonitrile (0.1% TFA), and then applied to a C-18 reverse phase HPLC column. Purified morphogen analog is harvested from the HPLC column.

The enhanced solubility of the resulting morphogen analog is measured by comparing the partition coefficient of the Ala 63-> Ser 63 mutein versus wild type hOP-1 dimer. It is contemplated that the Ala 63-> Ser 63 mutein has a higher solubility than native hOP-1. It is contemplated that, additional muteins having multiple hydrophobic to hydrophilic substitutions can be produced and characterized using the protocols described in this Example. The biological activity of the resulting morphogen analogs can be determined using one or more of the OP-1 activity assays described Example 1.

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# Example 3. Biological Activity of Finger 1. Finger 2, and Heel Peptides

The hOP-1-based peptides described in this example were produced and characterized prior to determination of the three-dimensional structure of hOP-1. These peptides either agonize or antagonize the biological activity of hOP-1. It is contemplated that, further refinements based upon the hOP-1 crystal structure, for example, the choice of more suitable sites for cyclizing peptides which constrain the peptide into a conformation that more closely mimics the shape of the corresponding region in hOP-1, may be used to further enhance the agonostic or antagonistic properties of such hOP-1-based peptides.

All of the peptides used in the following experiments, as well as their relationships with the mature hOP-1 amino acid sequence, are shown in Fig. 12. The finger 1-based peptides are designated F1 - 2; the heel-based peptides are designated H-1, H-n2 and H-c2; and the finger 2-based peptides are designated F2-2, and F2-3. Potential intra-peptide disulfide linkages are shown for each peptide. All the peptides were synthesized on a standard peptide synthesizer in accordance with the manufacturer's instructions. The peptides were deprotected, cyclized by oxidation, and then cleaved from resin prior to use.

In a first series of experiments, increasing concentrations of peptides F2-2 (Fig. 13A), F2-3 (Fig. 13B), Hn-2 (Fig. 13C) and Hc-2(Fig. 13D) were added to ROS cells either alone (open bars) or in combination with 40ng/ml soluble OP-1 (filled bars) and their effect on alkaline phosphatase activity measured. Soluble OP-1 is the form of OP-1 in which the pro-domain is still attached to the mature portion of OP-1 (see WO94/03600). A basal alkaline phosphatase activity is shown by the line and represents the alkaline phosphatase activity of cells incubated in the absence of both soluble OP-1 and peptide.

In Fig. 13A, peptide F2-2 at a concentration of about  $60~\mu\text{M}$  appears to double the basal alkaline phosphatase level and, in the presence of soluble OP-1, increases alkaline phosphatase activity by about 20% relative to soluble OP-1 alone. In Fig. 13B, peptide F2-3 at a concentration of about 0.01  $\mu\text{M}$  appears to increase the basal alkaline phosphatase level and, in the presence of soluble OP-1, increases alkaline phosphatase activity by about 20% relative to

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soluble OP-1 alone. Accordingly, both peptides F2-2 and F2-3, in the alkaline phosphatase assay, appear to act as weak OP-1 agonists.

In Fig. 13C, peptide H-n2 displays little or no effect on alkaline phosphatase activity either alone or in combination with soluble OP-1. Fig. 13D, peptide H-c2, at concentrations greater than about 5  $\mu$ M, appears to antagonize the activity of soluble OP-1.

In a second series of experiments, the ability of unlabeled soluble OP-1 and unlabeled peptides F1-2, F2-2, F2-3, H-n2 and H-c2 to displace <sup>125</sup>I labeled soluble OP-1 from ROS cell membranes was measured. The activities of peptides F2-2 and F2-3 relative to soluble OP-1 are shown in Fig. 14A, and the activities of peptides F1-2, H-n2 and H-c2 relative to soluble OP-1 are shown in Fig. 14B. OP-1 receptor-enriched plasma membranes of ROS cells were incubated for 20 hrs at 4°C with <sup>125</sup> I-labeled soluble OP-1 and unlabeled peptide. Receptor bound material was separated from unbound material by centrifugation at 39,500 x g. The resulting pellet was harvested and washed with 50mM HEPES buffer, pH7.4 containing 5mM MgCl<sub>2</sub> and 1mM CaCl<sub>2</sub> Radioactivity remaining in the pellet was determined by means of a gamma counter.

In Fig. 14A, peptide F2-2 (filled circles) soluble competes with soluble OP-1 with an Effective  $Dose_{50}$  ( $ED_{50}$ ) of about 1  $\mu$ M, but cannot completely displace soluble OP-1  $ED_{50}$  is the concentration of peptide to produce half maximal displacement of labeled soluble OP-1. Peptide F2-3 (filled triangles) competes and is able to completely displace soluble OP-1 with an  $ED_{50}$  of about 5  $\mu$ M. In Fig. 14B, peptide F1-2 (filled boxes), peptide H-n2 (open diamonds) and peptide H-c2 (open circles) all appear to exhibit little or no ability to displace iodinated soluble OP-1 from ROS cell membranes.

Although the peptide experiments appear promising, it is contemplated that resolution of the hOP-1 structure will enable the skilled practitioner to design constrained peptides that more closely mimic the receptor binding domains of human OP-1 and which are more effective at agonizing or antagonizing an hOP-1 mediated biological effect.

### Example 4. Elimination of a Binding Site on the Surface of OP-1

 $\alpha$ -2 macroglobulin, a protease scavenging protein known to bind proteins in serum and target them to the kidney for clearance from the body, binds OP-1. As described herein,  $\alpha$ -2's interaction sites on the OP-1 protein have been mapped. Accordingly, using the database and

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structural information provided herein, one can design an analog of OP-1 which eliminates one or more  $\alpha$ -2 macroglobulin interaction sites and provide an analog having enhanced bioavailability in the body. This same strategy can be applied for identifying and/or eliminating interaction sites for other binding proteins on the OP-1 surface.

## A. Identifying α-2 macroglobulin Binding Sites

OP-1 was determined to interact specifically with  $\alpha$ -2 macroglobulin in a standard competition binding assay, using immobilized, commercially available  $\alpha$ -2 macroglobulin, and labeled and unlabeled OP-1 protein. Truncated mature OP-1, wherein the first 22 amino acids have been cleaved from the mature form of OP-1 in a standard trypsin digest, bound  $\alpha$ -2 with 10-fold less affinity, indicating that the N terminal portion of the mature protein is involved in binding. This N-terminal portion of the protein, which is not part of the crystal structure, is positively charged and likely is highly flexible in solution. Elimination of this sequence does not interfere with OP-1 activity. Two cyclized peptides to all or a portion of the heel region, H-n2 and H1 (Cys<sub>71</sub> - Pro <sub>102</sub>, where Pro <sub>102</sub> has been changed to a cysteine to allow a disulfide bond between the two cysteines) also compete for binding; while peptides to the finger regions (F2-2, F2-3) do not compete.

 $\alpha$ -2 macroglobulin was determined not to interfere with OP-1's ability to stimulate alkaline phosphatase activity in a ROS cell assay. Accordingly,  $\alpha$ -2 macroglobulin binding does not appear to sterically inhibit OP-1 receptor binding.

### B. Design of Modified OP-1 Analog

The precise  $\alpha$ -2 macroglobulin interaction sites on OP-1 now can be mapped and an analog designed using the structure information provided herein. For example, the exact contact residues can be identified by creating model peptides like H-N2 and/or H1 in conjunction with an "alanine scan" mutagenesis program, wherein each residue is individually changed to an alanine in turn, and the constructs then tested for their ability to compete for binding. Once the contact residues are mapped, an analog can be designed which eliminates the contact residues without altering the overall structure of the heel region. Specifically, a template of the region can be called up on the computer from the database, and candidate replacement residues tested. The information in Table 8 identifies particularly useful candidate residues in the heel region which

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are solvent accessible, which likely are not available as epitopes and make good candidates for modification.

#### Equivalents.

The invention may be embodied in other specific forms without departing from the spirit or essential characteristics thereof. The foregoing embodiments are therefore to be considered in all respects illustrative rather than limiting on the invention described herein. Scope of the invention is thus indicated by the appended claims rather than by the foregoing description, and all changes that come within the meaning and range of equivalency of the claims are intended to be embraced therein.

#### SEQUENCE LISTING

#### (1) GENERAL INFORMATION:

- (i) APPLICANT: KECK, PETER GRIFFITH, DIANA L CARLSON, WILLIAM D RUEGER, DAVID C SAMPATH, KUBER T
- (ii) TITLE OF INVENTION: METHODS AND COMPOSITIONS FOR PRODUCING MORPHOGEN ANALOGS
- (iii) NUMBER OF SEQUENCES: 8
- (iv) CORRESPONDENCE ADDRESS:
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  - (E) COUNTRY: USA
  - (F) ZIP: 01748
- (v) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE: Floppy disk

  - (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
  - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
- (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER: US
    (B) FILING DATE:

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    - (A) LENGTH: 102 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (ix) FEATURE:
    - (A) NAME/KEY: Protein
    - (B) LOCATION: 1..102
    - (D) OTHER INFORMATION: /product= "hOP-1"
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

Cys Lys Lys His Glu Leu Tyr Val Ser Phe Arg Asp Leu Gly Trp Gln

5 10 15

Asp Trp Ile Ile Ala Pro Glu Gly Tyr Ala Ala Tyr Tyr Cys Glu Gly
20 25 30

Glu Cys Ala Phe Pro Leu Asn Ser Tyr Met Asn Ala Thr Asn His Ala 35 40 45

Ile Val Gln Thr Leu Val His Phe Ile Asn Pro Glu Thr Val Pro Lys 50 55 60

Pro Cys Cys Ala Pro Thr Gln Leu Asn Ala Ile Ser Val Leu Tyr Phe 65 70 75 80

Asp Asp Ser Ser Asn Val Ile Leu Lys Lys Tyr Arg Asn Met Val Val 85 90 95

Arg Ala Cys Gly Cys His 100

#### (2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 15 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (ix) FEATURE:
  - (A) NAME/KEY: Peptide
  - (B) LOCATION: 1..15
  - (D) OTHER INFORMATION: /product = "PEPTIDE F1-2"
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Cys Phe Arg Asp Leu Gly Trp Gln Asp Trp Ile Ile Ala Pro Cys
1 5 10 15

#### (2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 32 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (ix) FEATURE:
  - (A) NAME/KEY: Peptide
  - (B) LOCATION: 1..32
  - (D) OTHER INFORMATION: /product= "PEPTIDE H-1"
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Cys Ala Phe Pro Leu Asn Ser Tyr Met Asn Ala Thr Asn His Ala Ile 1 10 15 Val Gln Thr Leu Val His Phe Ile Asn Pro Glu Thr Val Pro Lys Cys 20 25 30

- (2) INFORMATION FOR SEQ ID NO:4:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:
    - (A) NAME/KEY: Peptide
    - (B) LOCATION: 1..13
    - (D) OTHER INFORMATION: /product= "PEPTIDE H-N2"
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Cys Leu Asn Ser Tyr Met Asn Ala Thr Asn His Ala Cys
1 10

- (2) INFORMATION FOR SEQ ID NO:5:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 11 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:
    - (A) NAME/KEY: Peptide
    - (B) LOCATION: 1..11
    - (D) OTHER INFORMATION: /product= "PEPTIDE H-C2"
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Cys Cys Phe Ile Asn Pro Glu Thr Val Cys Cys
1 10

- (2) INFORMATION FOR SEQ ID NO:6:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 11 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:
    - (A) NAME/KEY: Peptide
    - (B) LOCATION: 1..11
    - (D) OTHER INFORMATION: /product= "PEPTIDE F2-2"
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

- 74 -

Cys Tyr Phe Asp Asp Ser Ser Asn Val Ile Cys 1 5 10

- (2) INFORMATION FOR SEQ ID NO:7:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (ix) FEATURE:
    - (A) NAME/KEY: Peptide
    - (B) LOCATION: 1.16
    - (D) OTHER INFORMATION: /product = "PEPTIDE F2-3"
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Cys Tyr Phe Asp Asp Ser Ser Asn Val Ile Cys Lys Lys Tyr Arg Ser 1 5 10 15

- (2) INFORMATION FOR SEQ ID NO:8:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 98 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - · (ix) FEATURE:
    - (A) NAME/KEY: Protein
    - (B) LOCATION: 1..98
    - (D) OTHER INFORMATION: /product = "TGFB2"
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Cys Cys Leu Arg Pro Leu Tyr Ile Asp Phe Lys Arg Asp Leu Gly Trp
1 10 15

Lys Trp Ile His Glu Pro Lys Gly Tyr Asn Ala Asn Phe Cys Ala Gly
20 25 30

Ala Cys Pro Tyr Leu Trp Ser Ser Asp Thr Gln His Ser Arg Val Leu 35 40 45

Ser Leu Tyr Asn Thr Ile Asn Pro Glu Ala Ser Ala Ser Pro Cys Cys 50 60

Val Ser Gln Asp Leu Glu Pro Leu Thr Ile Leu Tyr Tyr Ile Gly Lys 65 70 75 80

Thr Pro Lys Ile Glu Gln Leu Ser Asn Met Ile Val Lys Ser Cys Lys 85 90 95

Cys Ser

#### **CLAIMS**

#### What is claimed is:

- 1 1. A computer system comprising:
- 2 (a) a memory having disposed therein atomic X-ray crystallographic co-ordinates
- defining at least a portion of human OP-1; and
- 4 (b) a processor in electrical communication with the memory; the processor comprising a
- 5 process which generates a molecular model having a three-dimensional shape representative of at
- 6 least a portion of human OP-1.
- 1 2. The system of claim 1, wherein the processor further comprises a process which
- 2 generates the molecular model having a solvent accessible surface representative of at least a
- 3 portion of human OP-1.
- 1 3. The system of claim 1, wherein said co-ordinates are stored on a computer readable
- 2 diskette.
- 1 4. The system of claim 1, wherein the molecular model is representative of at least a portion
- 2 of human OP-1 finger 1 region.
- 1 5. The system of claim 1 or 4, wherein the molecular model is representative of at least a
- 2 portion of the human OP-1 heel region.
- 1 6. The system of claim 1 or 4, wherein the molecular model is representative of at least a
- 2 portion of the human OP-1 finger 2 region.
- 1 7. The system of claim 6, wherein the molecular model is representative of at least a portion
- 2 of the human OP-1 heel region.
- 1 8. The system of claim 1, wherein the processor further identifies a morphogenic analog
- 2 having a three-dimensional shape and a solvent accessible surface corresponding to at least a
- 3 portion of the three-dimensional shape and the solvent accessible surface of human OP-1.

- 1 9. The system of claim 1, wherein the processor further identifies at least one candidate
- 2 amino acid defined by the co-ordinates, which upon modification enhances water solubility or
- 3 stability of human OP-1.
- 1 10. A method of producing a morphogenic analog having osteogenic protein-1 (OP-1) like
- 2 biological activity, the method comprising the steps of:
- 3 (a) providing a molecular model defining a three dimensional shape representative of at
- 4 least a portion of human OP-1;
- 5 (b) identifying a candidate analog having a three dimensional shape corresponding to the
- 6 three dimensional shape representative of at least a portion of human OP-1; and
- 7 (c) producing the candidate analog identified in step (b).
- 1 11. The method of claim 10, further comprising the step of determining whether the
- 2 compound produced in step (c) has an OP-1-like biological activity.
- 1 12. The method of claim 10, wherein the molecular model provided in step (a) is
- 2 representative of at least a portion of a finger 1 region of human OP-1.
- 1 13. The method of claim 10 or 12, wherein the molecular model provided in step (a) is
- 2 representative of at least a portion of a heel region of human OP-1.
- 1 14. The method of claim 10 or 12, wherein the model provided in step (a) is representative of
- 2 at least a portion of a finger 2 region of human OP-1.
- 1 15. The method of claim 14, wherein the molecular model provided in step (a) is
- 2 representative of at least a portion of a heel region of human OP-1.
- 1 16. The method of claim 10, wherein the analog comprises a plurality of charged moieties
- 2 spaced about the solvent accessible surface thereof and disposed in a spaced-apart relation
- 3 corresponding to charged moieties spaced about a portion of the solvent accessible surface of
- 4 human OP-1.

- 1 17. The method of claim 10, wherein steps (a) and (b) are performed by means of an
- 2 electronic processor.
- 1 18. The method of claim 17, wherein step (a) comprises storing a representation of at least a
- 2 portion of the atomic co-ordinates of human OP-1 in a computer memory.
- 1 19. A method of producing a morphogen analog that modulates an osteogenic protein-1 (OP-
- 2 1) mediated biological effect, the method comprising the steps of:
- 3 (a) providing in a computer memory atomic X-ray crystallographic co-ordinates defining
- 4 at least a portion of human OP-1;
- 5 (b) generating with a processor a molecular model having a three-dimensional shape and
- a solvent accessible surface representative of at least a portion of human OP-1,
- 7 (c) identifying a candidate morphogen analog having a three-dimensional structure shape
- 8 and a solvent accessible surface corresponding to the three-dimensional shape and the solvent
- 9 accessible surface of at least a portion of human OP-1;
- 10 (d) producing the candidate morphogen analog identified in step (c); and
- (e) determining whether the candidate morphogen analog produced in step (d) modulates
- 12 the OP-1 mediated biological effect.
- 1 20. The method of claim 11 or 19, further comprising the additional step of producing the
- 2 compound in a commercially useful quantity.
- 1 21. The method of claim 11 or 19, wherein said compound is a peptide.
- 1 22. A compound that modulates an OP-1 mediated biological effect produced by the method
- 2 of claim 11 or 19.
- 1 23. The compound of claim 22, wherein said compound agonizes the biological activity of
- 2 human OP-1.

FIG. 1A

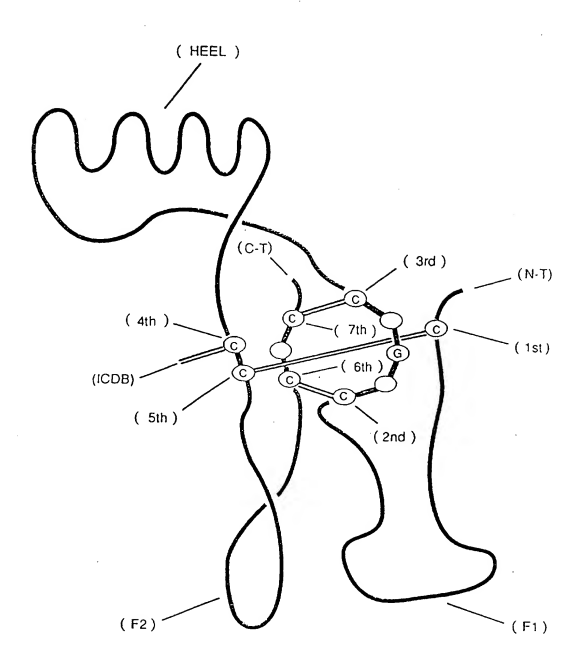


FIG. 1B

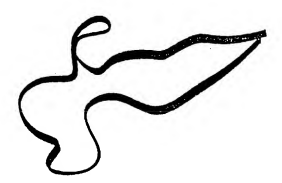


FIG. 1C

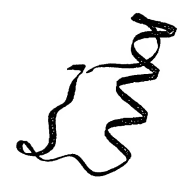
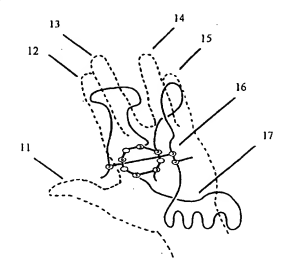


FIG. 1D



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### FIG. 1E



## FIG. 1F

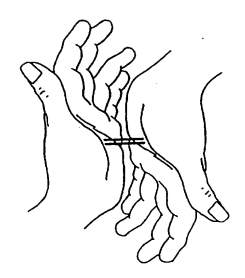
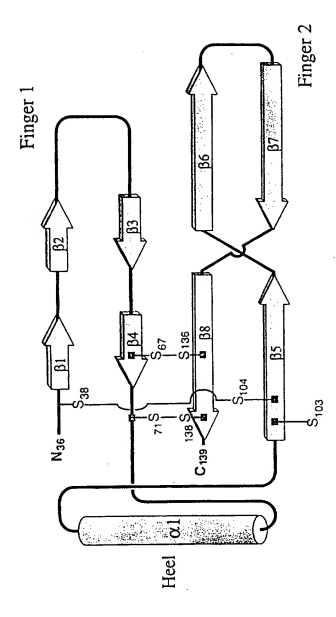


FIG. 2



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	FINGER 1				HEEL					FINGER 2		
2 °0	AYYCEGEC	N A N F C A G A C		100	<b>V</b> P C C	NPEASASPCC			<b>0</b> >	N.M.V.R.A.C.G.C.H.	ODLEPLTILYYIG - KTPK1EQLS'NMI'VKSCKCS	110
o> 9	ELYVSFR. DLGWQDWIIAPEGYAAYYC	>	4 0	-	LYHFINPET	A = A = A = A	0 2		- 3	I V I L K K Y R	PK-EQLS	100
o> s	LGWQDWI	K R D L G W · K W I H E P K G	3 0	o> 6	A VOTE	SRVCSEV	0 9		1 2 0	YFDDSSN	Y Y I G - K T	0
w	YVSFR.D	PLYIDFKRD		٥> 8	YMNATA			-	o>	LNAISVL	LEPLTIL	6
4 0>	CKKHEL	CCLRPL	2 0		AFPLNS	PYLWS	5 0		<del>-</del>	A P T Q	V · · S Q D L E P	8 0
	0P-1	TGF- 12			0P.1	TGF-N2				0P-1	TGF-p2	

Fig

FIG. 4A

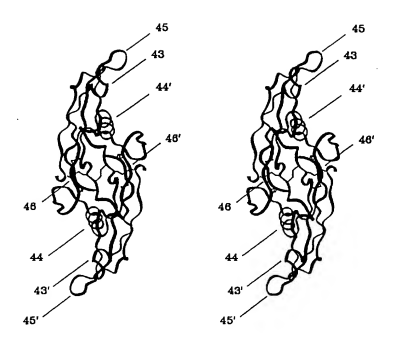


FIG. 4B

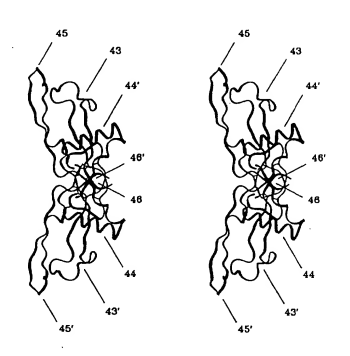
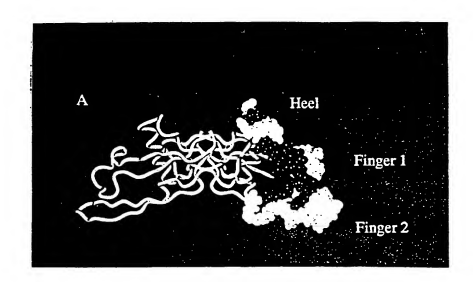


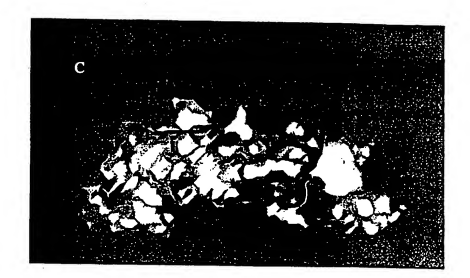
Fig. 5A



8/98 **Fig. 5B** 



Fig. 5C



	•		
£Ą-₹ÐT		Family	100
เฤ-ฯอา	10/98	<b>u.</b>	100 74
ह्य-म्बर		TGF-8	
4 <b>∂-</b> 4∂T		بارة	100 79 86 74 68
Othididal		Family 100	33 34 33
80nididn1	•	Activin 30 53 100 24 53	33 33 33
Agnididni		400 A 42	36 37 38
NODAL		36 37	43882
BMP-3		33 33 33 34 44	29 29 33 33
01-4 <b>0</b> 5	Fa mi 100 83	34 93	28 28 30 31
гсвем	BMP/OP Family 100 41 100 40 31 100 8	35 27 30	26 25 31 29
GDF-1	100 41 42	37 34 39	31 28 29 33 29
GDF-3	100 51 4 4 2 4 4 2 4 2 4 2 4 2 4 4 4 4 4 4 4	35 4 4 4 5 35	32 33 34 30 30 30 30 30 30 30 30 30 30 30 30 30
01-9MB	00 4 4 8 3 3 9 4 7 5 6 6 6 9 6 9 9 9 9 9 9 9 9 9 9 9 9 9 9	38 37 35	35 35 36 37
рокзегіи	100 66 68 43 37 37	36 34 32	30 31 33 33
6-4MB	100 79 63 63 64 64 65 78 78 78 78 78 78 78 78 78 78 78 78 78	39 35 35 34	33 30 30 30
6DF-5	100 50 53 53 4 7 7 4 4 3 3 9 4 4 3 4 4 3 4 4 3 4 4 4 4 4 4	35 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	33 33 34 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4
CDMP-2	001 8 5 5 4 5 8 5 4 7 6 7 6 7 6 7 6 7 6 7 6 7 6 7 6 7 6 7	42 40 36 36	34 2 3 3 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4
₹-4 <b>0</b> 5	100 82 80 80 80 80 80 80 80 84 47 47 47 47 47 47 47 47 47 47 47 47 47	38 38 39	35 35 35 35
6DF-6		44 40 37 38	35 35 35
<b>1-</b> 6√	100 50 51 51 51 50 50 50 50 50 50 50 50 50 50 50 50 50	42 38 34 42	35 33 37 37
ddp	100 4 4 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	38 37 34	36 36 36 36
BMP-4	76 76 76 77 76 76 76 76 76 76 76 76 76 7	44 42 44 37	33 34 44
S-9MB	100 74 4 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	4 4 4 8 6 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	35 35 35 35
ИМІМИ	100 64 65 65 65 65 65 65 65 65 65 65 65 65 65	49 41 42 37	33 33 34
A09	001 54 54 54 54 54 54 54 54 54 54	38 38 35 35	37 37 38 40 40
6-90	001 55 58 58 59 59 59 59 59 59 59 59 59 59 59 59 59	38 39 36 37	35 37 37 33
2-90	100 77 77 61 61 61 61 61 61 61 61 61 61 61 61 61	45 41 37 35	34 34 34 34 34 34 34 34 34 34 34 34 34 3
9-4MB	100 75 75 71 71 71 71 75 75 75 75 75 75 75 75 75 75 75 75 75	44 38 38 8	38 38 37
e-9MB	001 947 74 74 74 74 74 74 74 74 74 74 74 74 74 7	04 04 08 08 08 08	36 35 36 36
1-90	400 67 67 67 67 67 67 68 68 69 69 69 69 64 64 64 64 64 64 64 64 64 64 64 64 64	38 39 39	38 37 35 37
	+ OP41 + BMP-5 + BMP-6 OP-2 OP-3 + 60A UNIVIN + BMP-2 + BMP-2 + App Vg-1 GDF-6 GDF-5 GDF-5 BMP-10 GDF-3 GDF-3 GDF-3 GDF-3 GDF-3 GDF-10 GDF-10 GDF-10	NODAL InhibinßA InhibinßB InhibinßC	TGF-134 TGF-135 TGF-131 TGF-133

Fig. 6

PCT/US97/01071

ပ LKKYRNMVVRACGC APEGYAAYYCEGE 1 0 0> Ш MNATNHA I VQT L VHT APTOLNAISVLYFDDSSNV VSFRDLGWODW 5 0 > Ο> ∞ の対と田田口 FPL 4 0 > Fig. 7.4

Fig. 8.1

		Monomer	Dimer	Hidden Epitope	Surface	Modifiable
Re	sidue	% Area	% Area	% Area Residues	Modifiable	o Improve Solubility
36	GLN	71.89	71.89	0.00		
37	ALA	52.51	52.51	0.00		
39	LYS	62.19	62.19	0.00 EPITOPE		
40	LYS	39.26	39.26	0.00 EPITOPE		
41	HIS	27.13	27.13	0.00	•	
42	GLU	79.09	79.09	0.00 EPITOPE		
43	LEU	51.26	11.83	-39.43		
44	TYR	50.51	50.51	0.00 EPITOPE		
45	VAL	15.22	0.51	-14.71		
46	SER	23.02	23.02	0.00	*	
47	PHE	3.26	3.26	0.00		
48	ARG	76.89	76.89	0.00 EPITOPE		
49	ASP	68.71	52.15	-16.56 EPITOPE		
50	LEU	37.77	0.00	-37.77		
51	GLY	0.00	0.00	0.00		
52	TRP	40.99	34.53	-6.46	•	•
53	GLN	54.47	54.47	0.00 EPITOPE		
54	ASP	54.22	54.22	0.00 EPITOPE		
55	TRP	62.99	62.99	0.00 EPITOPE		
56	ILE	9.68	9.68	0.00	•	y
57	ILE	33.58	33.58	0.00 EPITOPE		
58	ALA	0.00	0.00	0.00		
59	PRO	34.01	34.01	0.00 EPITOPE		
60	GLU	60.90	60.90	0.00 EPITOPE		
61	GLY	0.00	0.00	0.00		
62	TYR	8.93	2.09	-6.84		
63	ALA	39.31	39.31	0.00	•	•

Fig. 8.2

64	ALA	14.78	0.00	-14.78		
65	TYR	26.22	26.22	0.00	•	*
66	TYR	48.32	15.41	-32.91		
67	CYS	1.67	1.67	0.00		
68	GLU	59.70	43.27	-16.43 EPITOPE		
69	GLY	0.00	0.00	0.00		
70	GLU	35.82	35.82	0.00 EPITOPE		
71	CYS	0.00	0.00	0.00		
72	ALA	43.27	43.27	0.00	•	•
73	PHE	39.54	39.54	0.00 EPITOPE		
74	PRO	96.68	96.68	0.00 EPITOPE		
75	LEU	1.72	1.72	0.00		
76	ASN	60.54	60.54	0.00 EPITOPE		
77	SER	73.24	73.24	0.00 EPITOPE		
78	TYR	104.34	104.34	0.00 EPITOPE		
79	MET	12.40	12.40	0.00		
80	ASN	46.31	46.31	0.00	*	
81	ALA	32.45	32.45	0.00	*	*
82	THR	34.63	5.99	-28.64		
83	ASN	84.54	38.00	-46.54		
84	HIS	71.01	0.26	-70.75		
85	ALA	0.00	0.00	0.00		
86	ILE	46.99	46.93	-0.06	•	*
87	VAL	64.29	1.95	-62.34		
88	GLN	18.05	4.31	-13.74		
89	THR	4.29	4.29	0.00		
90	LEU	50.95	29.43	-21.52		
91	VAL	39.39	8.51	-30.88		
92	HIS	26.42	26.42	0.00	*	

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Fig. 8.3

93	PHE	73.77	73.77	0.00 EPITOPE	
94	ILE .	57.23	32.03	-25.20 EPITOPE	
95	ASN	43.23	43.23	0.00 EPITOPE	
96	PRO	66.64	66.64	0.00 EPITOPE	
97	GLU	88.25	88.25	0.00 EPITOPE	
98	THR	52.59	48.71	-3.88 EPITOPE	
99	VAL	25.83	0.00	-25.83	
100	PRO	89.22	30.78	-58.44	
101	LYS	35.15	35.15	0.00	*
102	PRO	0.00	0.00	0.00	
103	CYS	79.14	27.13	-52.01	
104	CYS	5.39	5.39	0.00	
105	ALA	44.46	5.15	-39.31	
106	PRO	11.24	2.30	-8.94	
107	THR	21.76	21.76	0.00	•
108	GLN	53.40	53.40	0.00 EPITOPE	
109	LEU	29.98	7.79	-22.19	
110	ASN	35.00	35.00	0.00	*
111	ALA	23.61	23.61	0.00	•
112	ILE	22.72	22.72	0.00	•
113	SER	38.55	38.55	0.00	. •
114	VAL	1.15	1.15	0.00	
115	LEU	36.05	36.05	0.00 EPITOPE	
116	TYR	18.62	18.62	0.00	
117	PHE	46.55	46.55	0.00 EPITOPE	
118	ASP	32.53	32.53	0.00 EPITOPE	
119	ASP	84.02	84.02	0.00 EPITOPE	
120	SER	48.35	48.35	0.00 EPITOPE	
121	SER	68.39	68.39	0.00 EPITOPE	

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Fig. 8.4

122	ASN	63.15	63.15	0.00 EPITOPE		
123	VAL	41.27	41.27	0.00 EPITOPE		
124	ILE	34.51	34.51	0.00 EPITOPE		
125	LEU	63.34	63.34	0.00 EPITOPE		
126	LYS	54.81	54.81	0.00 EPITOPE		
127	LYS	48.78	48.78	0.00 EPITOPE		
128	TYR	34.23	32.55	-1.68	•	*
129	ARG	63.25	62.85	-0.40 EPITOPE		
130	ASN	62.31	40.62	-21.69		
131	MET	32.35	7.44	-24.91		
132	VAL	16.38	16.38	0.00		
133	VAL	7.50	0.07	-7.43		
134	ARG	65.10	65.10	0.00		
135	ALA	47.10	47.10	0.00	•	*
136	CYS	0.29	0.29	0.00		
137	GLY	0.00	0.00	0.00		
138	CYS	0.00	0.00	0.00		
139	HIS	47.68	18.94	-28.74		

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Fig. 9

Ridge R	esidues	Receptor Sites			
B90	Leu	Heel			
B91	Val	Heel			
B92	His	Heel			
B93	Phe	Heel	•		
B94	lle	Heel	ł		
B95	Asn	Heel			
B96	Pro	Heel			
B97	Glu	Heel	• 1		
B98	Thr	Heel	<u>.</u>		
A48	Arg	Finger 1	*		
A49	Asp	Finger 1	i		
A50	Leu	Finger 1			
A51	Gly	Finger 1			
A52	Trp	Finger 1	i		
A53	GIn	Finger 1	•		
A54	Asp	Finger 1			
A55	Trp	Finger 1			
A56	lle	Finger 1			
A57	lle	Finger 1			
A58	Ala	Finger 1			
A59	Pro	Finger 1			
A60	Glu	Finger 1	*		
A116	Tyr	Finger 2			
A117	Phe	Finger 2	•		
A118	Asp	Finger 2			
A119	Asp	Finger 2	*		
A120	Ser	Finger 2	•		
A121	Ser	Finger 2	•		
A122	Asn	Finger 2			
A123	Val	Finger 2	i		
A124	lle	Finger 2	3		
A125	Leu	Finger 2	I I		
A126	Lys	Finger 2			
A127	Lys	Finger 2	1		
A128	Tyr	Finger 2			
A129	Arg	Finger 2	*		

Fig. 10

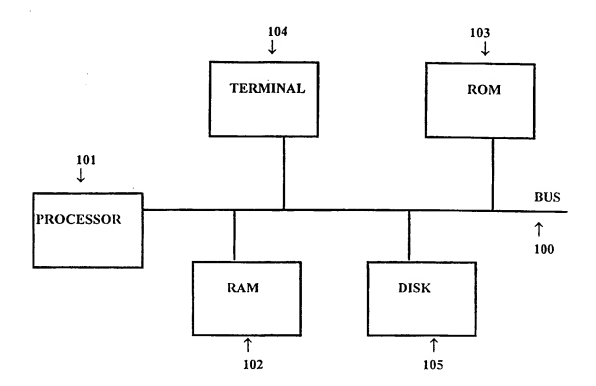


Fig. 11A

Residue in 1st Chain	Residue in 2nd Chain	Distance (A)
Ala-105	Ala-105	3.61
Cys-103	Cys-103	3.95
Asn-83	Asn-130	4.01
Thr-82	Asn-130	4.20

Fig. 11B

Residue in Finger 1	Residue in Finger 2	Distance (A)
Ala-58	Val-114	3.30
Tyr-65	Val-133	3.93
Ala-58	Leu-115	3.98
llc-57	Leu-115	4.62
lle-56	Tyr-116	4.54
Trp-55	Tyr-116	4.74

# FIGURE 12

OP-1:	CKKHELYVSFRDLGWQDWIIAPEGYAAYYCEGECAFPLNSYMNATNHAIVQTLVHFINPETVPKPCCAPTQLNAISVLYFDDSSNVILKKYRNMVVRACGCH   <finger 1<="" th=""></finger>
Finger 1 Peptides:	
F1-2:	CFRDLGWQDWIIAPC
Heel Peptides:	
1-н	CAFPLNSYMJATNHAIVQTLVHFINPETVPKC
Н-п2:	CLUSYMIATNHAC
й- <b>с</b> 2:	19/98
Finger 2 Peptides	
F2-2	CYFDDSSNVIC
F2-3	CYFDDSSNVICKKYRS

Fig. 13A

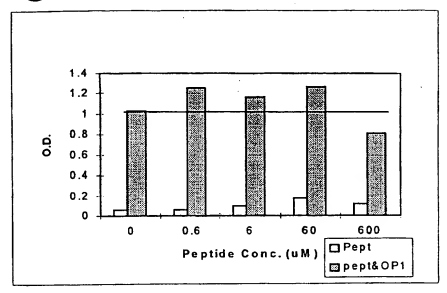


Fig. 13B

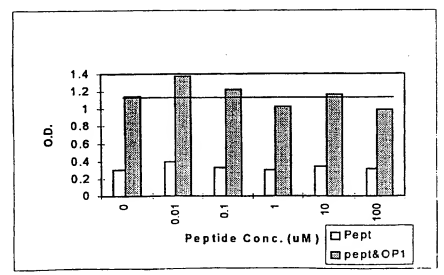


Fig. 13C

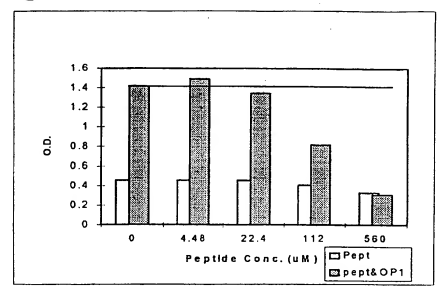
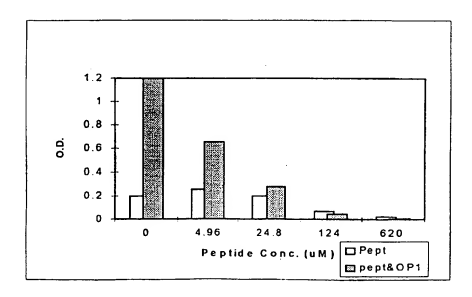


Fig. 13D



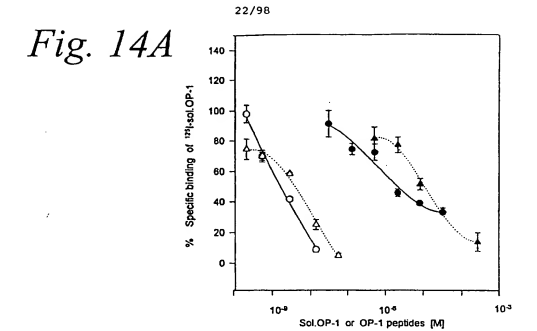


Fig. 14B

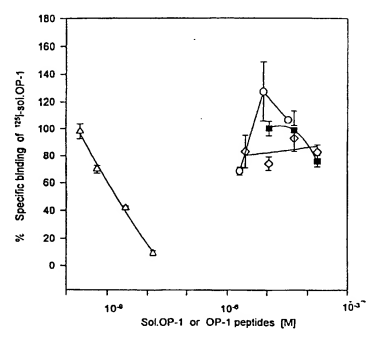


Fig. 15.1

A	tom	Resi	due	Chain	x	Y	z
1	N	GLN	36	A		-13.365	11.899
2	CA	GLN	36	A		-13.227	11.283
3	C	GLN	36	A		-12.220	10.237
4	0	GLN	36	A		-11.030	10.439
5	CB	GLN	36	A		-12.644	12.291
6	ÇG	GLN	36	A		-11.130	12.646
7	CD	GLN	36	λ		-10.667	13.351
8	OE1	GLN	36	A		-11.405	13.452
9	NE2	GIN	36	A	2.743	-9.471	13.880
10	N	ALA	37	A		-12.854	9.097
11	CA	ALA	37	A		-12.466	8.044
12	С	ALA	37	A		-11.639	7.174
13	0	ALA	37	A		-11.899	7.227
14	CB	ALA	37	A		-13.708	7.294
15	N	CYS	38	A		-10.694	6.369
16	CA	CYS	38	A	2.177	-9.971	5.682
17	С	CYS	38	A		-10.860	4.675
18	0	CYS	38	A		-11.266	3.753
19	CB	CYS	38	A	2.821	~8.791	5.057
20	SG	CYS	38	A	1.945	-8.415	3.597
21	N	LYS	39	A		-11.141	4.862
22	CA	LYS	39	A		-11.883	3.903
23	С	LYS	39	A		-11.277	3.660
24	0	LYS	39	A		-10.382	4.358
25	CB	LYS	39	A		-13.348	4.365
26	CG	LYS	39	A		-13.470	5.883
27	CD	LYS	39	λ		-14.824	6.505
28	CE	LYS	39	A		-14.585	8.025
29	NZ	LYS	39	A		-15.799	8.829
30	N	LYS	40	A		-11.748	2.658
31	CA	LYS	40	A		-11.299	2.397
32	С	LYS	40	A		-11.916	3.394
33	0	LYS	40	A		-12.991	3.851
34	CB	LYS	40	A		-11.743	1.058
35	CG	LYS	40	A		-11.024	0.591
36	CD	LYS	40	A		-11.965	-0.285
37	CE	LYS	40	λ		-12.585	-1.352
38	NZ	LYS	40	A		-13.536	-2.104
39	N	HIS	41	λ		-11.330	3.782
40	CA	HIS	41	λ		-11.957	4.717
41	С	HIS	41	A		-11.660	4.308
42	0	HIS	41	A		-10.901	3.384
43	CB	BIS	41	A		-11.406	6.101
44	CG	HIS	41	A		-11.606	6.758
45	ND1	HIS	41	A	-5.353	-12.221	7.907

Fig. 15.2

A1	tom	Residu	ie Ch	ain X	Y	<u>z</u>
46	CD2	HIS 4	41 A	-4.544	-10.869	6.408
47	CE1		41 A		-11.833	8.293
48	NE2		41 A		-11.017	7.380
49	NEZ N		42 A	•	-12.183	4.920
50	CA		42 A		-12.077	4.368
51	C		42 A		-10.923	4.856
	0		42 A		-10.417	5.925
52	СВ		42 A		-13.320	4.667
53			42 A		-14.387	3.691
54	CG		42 A		-15.373	3.342
55	CD		42 A		-16.555	3.062
56	OE1		42 A		-14.955	3.347
57	OE2		43 A		-10.414	4.163
58	N			-13.426		4.751
59	CA			-14.554		3.765
60	C			-14.352		2.600
61	0			-12.846		4.589
62	СВ		43 A	-12.886		5.690
63	CG		43 A	-12.880		5.031
64	CD1		43 A			
65	CD2	_	43 A	+13.968		6.695 4.196
66	N		44 A	-15.723 -16.879		3.365
67	CA		44 A			3.638
68	С		44 A	-17.493		4.790
69	0		44 A	-17.838		
70	CB		44 A		-10.903	3.800
71	CG	_	44 A		-11.049	2.820
72	CD1		44 A		-10.566	3.189
73	CD2		44 A		-11.580	1.592
74	CE1		44 A		-10.473	2.308
75	CE2		44 A		-11.489	0.699
76	CZ		44 A		-10.877	1.042
77	OH		44 A		-10.553	0.068
78	N		45 A			2.740
79	CA		45 A	-18.427		3.122
80	С		45 A	-19.847		2.610
81	0		45 A	-20,040		1.461
82	CB		45 A	-17.796		2.507
83	CG1		45 A	-18.457		2.960
84	CG2		45 A			2.939
85	N		46 A	-20.877		3.431
86	CA		46 A			3.059
87	С		46 A			2.918
88	0		46 A			3.707
. 89	CB	SER	46 A			
90	OG	SER	46 A	-24.417	7 -7.335	3.799

Fig. 15.3

A	Atom		duc	Chain	x	Υ	
91	N	PHE	47	A	-23.389	-4.851	1.705
92	CA	PHE	47	A	-23.751	-3.481	1.472
93	C	PHE	47	A	-24.740	-2.959	2.450
94	0	PHE	47	A	-24.979	-1.770	2.498
95	CB	PHE	47	A	-24.352	-3.321	0.142
96	CG	PHE	47	A	-23.337	-3.619	-0.910
97	CD1	PHE	47	A	-22.233	-2.845	-1.022
98	CD2	PHE	47	A	-23.634	-4.545	-1.868
99	CE1	PHE	47	A	-21.486	-2.920	-2.152
100	CE2	PHE	47	A	-22.889	-4.598	-3.018
101	CZ	PHE	47	A	-21.825	-3.765	-3.173
102	N	ARG	48	A	-25.371	-3.781	3.275
103	CA	ARG	48	A	-26.102	-3.279	4.414
104	С	ARG	48	A	-25.162	-2.527	5.278
105	0	ARG	48	A	-25.411	-1.399	5.572
106	CB	ARG	48	A	-26.719	-4.406	5.231
107	CG	ARG	48	A	-27.809	-5.147	4.442
108	CD	ARG	48	A	-28.661	-6.042	5.341
109	NE	ARG	48	A	-29.918	-6.386	4.698
110	CZ	ARG	48	A	-30.415	-7.627	4.795
111	NH1	ARG	48	A	-29.715	-8.597	5.456
112	NH2	ARG	48	A	-31.623	-7.961	4.241
113	N	ASP	49	A	-24.037	-3.036	5.731
114	CA	ASP	49	A	-23.280	-2.378	6.778
115	C	ASP	49	A	-22.753	-1.006	6.372
116	0	ASP	49	A	-22.313	-0.188	7.168
117	CB	ASP	49	A	-22.095	-3.224	7.190
118	CG	ASP	49	A	-22.486	-4.646	7.506
119	OD1	ASP	49	A	-21.605	-5.482	7.303
120	OD2	ASP	49	A	-23.619	-4.962	7.924
121	N	LEU	50	A	-22.770	-0.677	5.092
122	CA	LEU	50	A	-22.319	0.638	4.686
123	C	LEU	50	A	-23.510	1.477	4.402
124	0	LEU	50	A	-23.448	2.588	3.870
125	CB	LEU	50	A	-21.468	0.539	3.433
126	CG	LEU	50	A	-20.287	-0.404	3.582
127	CD1	LEU	50	Α	-20.061	-1.030	2.227
128	CD2	LEU	50	A	-19.052	0.319	4.121
129	N	GLY	51	A	-24.651	0.915	4.749
130	CA	GLY	51	A	-25.907	1.551	4.454
131	С	GLY	51	A	-26.245	1.497	2.985
132	0	GLY	51	A	-27.355	1.857	2.609
133	N	TRP	52	A	-25.389	1.072	2.054
134	CA	TRP	52	A	-25.828	1.011	0.660
135	С	TRP	52	A	-27.045	0.114	0.381

Fig. 15.4

Atom		Resi	Residue		x	<u>Y</u>	Z
136	0	TRP	52	A	-27.129	-0.483	-0.668
137	CB	TRP	52	A	-24.614	0.556	-0.189
138	CG	TRP	52	A	-23.484	1.602	-0.152
139	CD1	TRP	52	A	-23.615	2.896	0.315
140	CD2	TRP	52	A	-22.092	1.295	-0.610
141	NE1	TRP	52	A	-22.430	3.454	0.213
142	CE2	TRP	52	A	-21.489	2.641	-0.293
143	CE3	TRP	52	A	-21.262	0.304	-1.105
144	CZ2	TRP	52	A	<b>-20.133</b>	2.852	-0.442
145	CZ3	TRP	52	A	-19.913	0.575	-1.247
146	CH2	TRP	52	A	-19.360	1.810	-0.907
147	N	GLN	53	A	-28.072	-0.088	1.197
148	CA	GLN	53	A	-28.758	-1.377	1.098
149	C	GLN	53	A	-29.798	-1.458	0.072
150	0	GLN	53	, <b>A</b>	-30.460	-2.475	-0.062
151	CB	GLN	53	A	-29.504	-1.835	2.360
152	CG	GLN	53	A	-29.756	-0.759	3.403
153	CD	GLN	53	A	-31.038	-1.078	4.192
154	OE1	GLN	53	A	-31.953	-0.232	4.230
155	NE2	GLN	53	A	-31.163	-2.269	4.840
156	N	ASP	54	A	-29.992	-0.405	-0.689
157	CA	ASP	54	A	-31.357	0.003	-0.992
158	С	ASP	54	A	-31.136	0.729	-2.255
159	0	ASP	54	A	-31.000	1.942	-2.119
160	СВ	ASP	54	A	-31.873	0.982	0.073
161	CG	ASP	54	A	-30.722	1.807	0.710
162	OD1	ASP	54	A	-30.861	2.211	1.892
163	OD2	ASP	54	A	-29.698	2.012	0.027
164	N	TRP	55	A	-31.067	0.175	-3.470
165	CA	TRP	55	A	-30.322	0.915	-4.515
166	С	TRP	55	A	-29.770	-0.083	-5.462
167	0	TRP	55	A	-29.915	0.104	-6.662
168	CB	TRP	55	A	-29.011	1.599	-4.158
169	CG	TRP	55	A	-29.091	3.035	-3.787
170	CD1	TRP	55	A	-30.088	3.925	-4.101
171	CD2	TRP	55	A	-27.886	3.623	-3.134
172	NE1	TRP	55	A	-29.608	5.108	-3.722
173	CE2	TRP	55	A	-28.348	5.080	-3.173
174	CE3	TRP	55	A	-26.706	3.288	-2.489
175	CZ2	TRP	55	A	-27.508	6.047	-2.602
176	CZ3	TRP	55	A	-25.932	4.297	-1.921
177	CH2	TRP	55	A	-26.304	5.642	-1.988
178	N	ILE	56	A	-29.151	-1.108	-4.812
179	CA	ILE	56	A	-28.323	-2.142	-5.400
180	c	ILE	56	А	-29.209	-3.352	-5.699

Fig. 15.5

Atom		Residuc		Chain	X	Y	<u>z</u>
181	0	ILE	56	λ	-29.900	-3.795	-4.792
182	CB	ILE	56	A	-27.221	-2.452	-4.390
183	CG1	ILE	56	Ä	-25.998	-1.743	-4.815
184	CG2	ILE	56	Ä	-26.917	-3.924	-4.291
185	CD1	ILE	56	A	-25.964	-0.422	-4.122
186	N	ILE	57	A	-29.260	-3.924	-6.907
187	CA	ILE	57	A	-29.961	-5.182	-7.093
188	C	ILE	57	Ä	-28.927	-6.288	-6.809
189	0	ILE	57	A	-29.163	-7.475	-6.713
190	СВ	ILE	57	A	-30.481	-5.241	-8.549
191	CG1	ILE	57	A	-31.405	~4.079	-8.780
192	CG2	ILE	57	A	-31.273	-6.511	-8.844
193	CD1	ILE	57	A	-31.745		-10.274
194	N	ALA	58	A	-27.684	~5.914	-6.623
195	CA	ALA	58	A	-26.550	-6.030	-7.556
196	C	ALA	58	A	-25.789	-7.248	-7.200
197	ō	ALA	58	A	-26.272	-8.230	-7.747
198	СВ	ALA	58	A	-25.682	-4.813	-7.386
199	N	PRO	59	A	-24.748	-7.272	-6.411
200	CA	PRO	59	A	-24.570	-8.349	-5.438
201	C	PRO	59	A	-25.086	-7.952	-4.063
202	ō	PRO	59	A	-25.265	-6.789	-3.797
203	СВ	PRO	59	А	-23.068	-8.697	-5.439
204	CG	PRO	59	A	-22.447	-7.366	-5.730
205	CD	PRO	59	A	-23.408	-6.817	-6.757
206	N	GLU	60	A	-25.376	-8.795	-3.088
207	CA	GLU	60	A	-25.696	-8.242	-1.777
208	C	GLU	60	A	-24.492	~7.832	-0.980
209	0	GLU	60	A	-24.571	-7.233	0.078
210	СВ	GLU	60	· A	-26.502	-9.249	-0.927
211	CG	GLU	60	A	-25.753	-10.380	-0.180
212	CD	GLU	60	A	-24.990	-9.971	1.084
213	OE1	GLU	60	A	-25.319	-8.939	1.717
214	OE2	GLU	60	A	-24.089	-10.738	1.457
215	N	GLY	61	A	-23.307	-8.146	-1.448
216	CA	GLY	61	A	-22.099	-7.722	-0.784
217	С	GLY	61	A	-21.009	-8.349	-1.571
218	0	GLY	61	A	-21.289	-8.862	-2.648
219	N	TYR	62	A	-19.777	-8.343	-1.088
220	CA	TYR	62	A	-18.689	-8.872	-1.879
221	С	TYR	62	A	-17.524	-9.144	-0.981
222	0	TYR	62	A	-17.554	-8.747	0.169
223	СВ	TYR	62	A	-18.362	-7.838	-2.934
224	CG	TYR	62	A	-17.588	-6.667	-2.413
225	CD1	TYR	62	A	-18.181	-5.490	-1.956

## Fig. 15.6

Atom		Resid	Residue		X	Y	<u>z</u>
226	CD2	TYR	62	A	-16.236	-6.785	-2.513
227	CE1	TYR	62	A	-17.379	-4.412	-1.599
228	CE2	TYR	62	A	-15.447	-5.739	-2.169
229	cz	TYR	62	A	-16.004	-4.571	-1.715
230	ОН	TYR	62	A	-15.097	-3.565	-1.383
231	N	ALA	63	A	-16.476	-9.808	-1.456
232	CA	ALA	63	A	-15.231	-9.961	-0.740
233	C	ALA	63	А	-14.242	-8.820	-0.961
234	0	ALA	63	A	-13.659	-8.712	-2.023
235	СВ	ALA	63	A	-14.655	-11.248	-1.205
236	N	ALA	64	A	-14.020	-7.948	0.007
237	CA	ALA	64	A	-13.051	-6.893	-0.114
238	С	ALA	64	A	-11.606	-7.304	0.192
239	0	ALA	64	A	-10.608	-6.800	-0.298
240	CB	ALA	64	A	-13.454	-5.781	0.832
241	N	TYR	65	A	-11.469	-8.294	1.060
242	CA	TYR	65	A	-10.215	-8.675	1.672
243	С	TYR	65	A.	-9.503	-7.544	2.281
244	0	TYR	65	Α	-9.863	-6.400	2.147
245	СВ	TYR	65	A	-9.2 <b>9</b> 0	-9.333	0.678
246	CG	TYR	65	A	-9.846	-10.614	0.126
247	CD1	TYR	65	A	-10.300	-11.608	0.939
248	CD2	TYR	65	A	-9.927	-10.753	-1.219
249	CE1	TYR	65	A	-10.905	-12.716	0.405
250	CE2	TYR	65	A	-10.526	-11.847	-1.767
251	CZ	TYR	65	A		-12.799	-0.951
252	OH	TYR	65	A	-11.729	-13.859	-1.515
253	N	TYR	66	A	-8.459	-7.830	2.998
254	CA	TYR	66	A	-7.615	-6.771	3.442
255	С	TYR	66	A	-6.303	-7.401	3.710
256	0	TYR	66	A	-6.198	-8.619	3.836
257	CB	TYR	66	A	-8.154	-6.109	4.712
258	CG	TYR	66	A	-8.091	-6.906	5.984
259	CD1	TYR	66	A	-8.948	-7.941	6.159
260	CD2	TYR	66	A	-7.215	-6.546	6.978
261	CE1	TYR	66	A	-8.929	-8.634	7.333
262	CE2	TYR	66	A	-7.183	-7.227	8.155
263	CZ	TYR	66	A	-8.041	-8.273	8.310
264	OH	TYR	66	A	-8.024	-9.037	9.472
265	N	CYS	67	A	~5.286	-6.571	3.812
266	CA	CYS	67	A	-3.921	-7.055	3.949
267	C	CYS	67	A	-3.364	-6.825	5.358
268	. 0	CYS	67	A	-3.426	-5.705	5.888
269	CB	CYS	67	A	-2.971	-6.328	3.056
270	SG	CYS	67	A	-3.158	-6.344	1.310

Fig. 15.7

	Atom	Re	sidue	Chain	x	Y	z
271	N	GLU	68	A	-2,792	-7.828	6.031
272	CA	GLU	68	A	-2.322		7.318
273	С	GLU	68	A	-1.275		7.726
274	0	GLU	68	A	-1.438		7.481
275	CB	GLU	68	A	-3.408		8.334
276	CG	GLU	68	A	-3.092	-6.620	9.488
277	CD	GLU	68	A	-3.797	-7.134	10.693
278	OE1	GLU	68	A	-3.899	-6.397	11.674
279	OE2	GLU	68	A	-4.246	-8.282	10.661
280	N	GLY	69	A	-0.219	-7.857	8.348
281	CA	GLY	69	A	0.772	-8.701	8.979
282	C	GLY	69	A	2.107	-7.997	8.920
283	0	GLY	69	A	2.212	-6.844	8.501
284	N	GLU	70	A	3.192	-8.624	9.316
285	CA	GLU	70	A	4.349	-7.810	9.600
286	С	GLU	70	A	5.364	-7.795	8.496
287	0	GLU	70	A	5.603	-8.818	7.881
288	CB	GLU	70	A .	4.996	-8.309	10.865
289	CG	GLU	70	A	5.202	-7.080	11.735
290	CD	GLU	70	A	6.392	-7.339	12.618
291	OE1	GLU	70	A	7.092	-6.368	12.941
292	OE2	GLU	70	A	6.637	-8.516	12.968
293	N	CYS	71	A	5.988	-6.666	8.222
294	CA	CYS	71	A	7.049	-6.657	7.253
295	С	CYS	71	A	8.436	-6.637	7.832
296	0	CYS	71	A	8.961	-5.627	8.241
297	CB	CYS	71	A	6.787	-5.467	6.371
298	SG	CYS	71	A	5.598	-5.970	5.102
299	N	ALA	72	A	9.141	-7.740	7.914
300	CA	ALA	72	A	10.322	-7.789	8.729
301	C	ALA	72	A	11.166	-8.899	8.174
302	0	ALA	72	A	10.670	-9.710	7.406
303 304	CB	ALA	72	A	9.874	-8.066	10.139
305	N CA	PHE	73	A	12.457	-9.057	8.463
306	C	PHE PHE	73	A	13.195	-10.119	7.773
307	0	PHE	73 73	A		-11.435	8.328
308	СВ	PHE	73 73	A		-11.454	9.530
309	CG	PHE	73	A	14.723	-9.995	7.990
310	CD1	PHE	73	A	15.255	-8.809	7.230
311	CD2	PHE	73 73	A	15.243	-8.804	5.868
312	CE1	PHE	73 73	A	15.746	-7.737	7.919
313	CE2	PHE	73 73	A	15.709	-7.716	5.190
314	CZ	PHE	73 73	A A	16.225	-6.647	7.217
315	N	PRO	74		16.203	-6.634	5.857
-13	••	FRU	, 4	A	12.570	-12.531	7.695

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Fig.	<i>15.8</i>

A	tom	Resi	due	Chain	x	Y	<u>z</u>
316	CA	PRO	74	A	12.982	-12.802	6.339
317	С	PRO	74	A	11.990	-12.148	5.485
318	0	PRO	74	Α.	10.891	-12.643	5.423
319	CB	PRO	74	A	12.945	-14.284	6.176
320	CG	PRO	74	A	12.664	-14.772	7.587
321	CD	PRO	74	A	11.893	-13.668	8.292
322	N	LEU	75	A	12.254	-11.051	4.786
323	CA	LEU	75	A	11.328	-10.466	3.831
324	С	LEU	75	A	11.299	-11.508	2.762
325	0	LEU	75	A	11.992	-11.532	1.756
326	CB	LEU	75	A	11.912	-9.144	3.346
327	CG	LEU	75	A	10.922	-8.023	3.399
328	CD1	LEU	75	A	9.924	-8.232	4.514
329	CD2	LEU	75	A	11.680	-6.741	3.622
330	N	ASN	76	A	10.419	-12.442	3.038
331	CA	ASN	76	A	10.337	-13.665	2.288
332	С	asn	76	A	9.758	-13.535	0.863
333	0	asn	76	A	8.766	-14.166	0.569
334	CB	asn	76	A	9.548	-14.511	3.229
335	CG	nea	76	A	9.704	-15.945	2.935
336	OD1	asn	76	A	9.686	-16.766	3.862
337	ND2	asn	76	A	9.760	-16.324	1.643
338	N	SER	77	A	10.263	-12.752	-0.109
339	CA	SER	77	A	9.917	-12.843	-1.545
340	С	SER	77	A	8.445	-12.920	-1.915
341	0	SER	77	A		-11.897	-2.284
342	CB	SER	77	A	10.599	-14.059	-2.195
343	OG	SER	77	A	10.201	-15.273	-1.560
344	N	TYR	78	A	7.776	-14.081	-1.847
345	CA	TYR	78	A	6.343	-13.977	-1.994
346	С	TYR	78	A	5.734	-13.258	-0.792
347	0	TYR	78	A	4.827	-13.833	-0.216
348	СВ	TYR	78	A	5.683	-15:337	-2.079
349	CG ·	TYR	78	A	5.162	-15.723	-3.465
350	CD1	TYR	78	A	4.144	-16.681	-3.432
351	CD2	TYR	78	A	5.865	-15.473	-4.665
352	CE1	TYR	78	A	3.985	-17.578	-4.480
353	CE2	TYR	78	A	5.706	-16.377	-5.729
354	CZ	TYR	78	A	4.859	-17.489	-5.557
355	OH	TYR	78	A	4.965	-18.610	-6.363
356	N	MET	79	A	6.220	-12.058	-0.418
357	CA	MET	79	A	5.629	-11.035	0.425
358	С	MET	79	, <b>A</b>	5.931	-9.703	-0.270
359	0	MET	79	A	5.976	-8.606	0.269
360	CB	MET	79	A	6.271	-10.880	1.754

Fig. 15.9

A	tom	Resi	iduc	Chain	x	Y	z
361	CG	MET	79	A	6.288	-12.036	2.716
362	SD	MET	79	A		-11.247	4.193
363	CE	MET	79	A		-12.664	5.230
364	N	ASN	80	A	6,171	-9.795	-1.562
365	CA	ASN	80	A	6.444	-8.675	-2.417
366	С	ASN	80	A	7.324	-7.534	-1.973
367	0	ASN	80	A	7.169	-6.458	-2.499
368	CB	ASN	80	A	5.129	-8.059	-2.909
369	CG	ASN	80	A	5.403	-7.179	-4.135
370	OD1	ASN	80	A	5.383	-5.934	-4.172
371	ND2	ASN	80	A	5.792	-7.856	-5.212
372	N	ALA	81	A	8.295	-7.505	-1.068
373	CA	ALA	81	A	9.543	-6.854	-1.431
374	С	ALA	81	A	9.731	-5.440	-1.931
375	0	ALA	81	A	10.248	-4.546	-1.301
376	CB	ALA	81	A	10.249	-7.718	-2.461
377	N	THR	82	Α	9.352	-5.062	-3.108
378	CA	THR	82	A	10.122	-4.047	-3.818
379	С	THR	82	A	11.649	-4.060	-3.652
380	0	THR	82	A	12.323	-4.730	-4.441
381	CB	THR	82	A	9.752	-2.568	-3.534
382	OG1	THR	82	A	9.779	-2.312	-2.158
383	CG2	THR	82	A	8.458	-2.245	-4.220
384	N	asn	83	A	12.288	-3.380	-2.692
<b>38</b> 5	CA	ASN	83	λ	13.708	-3.070	-2.868
386	С	ASN	83	A	13.915	-1.957	-1.951
387	0	ASN	83	A	14.715	-1.956	-1.057
388	CB	ASN	83	A	14.082	-2.493	-4.184
389	CG	ASN	83	A	14.958	-3.390	-4.974
390	OD1	ASN	83	A	15.593	-2.957	-5.919
391	ND2	ASN	83	A	14.990	-4.682	-4.709
392	N	HIS	84	A	13.196	-0.870	-2.081
393	CA	HIS	84	A	13.067	0.094	-1.008
394	С	HIS	84	A	12.526	-0.632	0.181
395	0	HIS	84	A	12.864	-0.201	1.261
396	CB	HIS	84	A	12.095	1.187	-1.379
397	CG	HIS	84	A	12.020	2.264	-0.350
398	ND1	HIS	84	A	11.255	2.299	0.730
399	CD2	HIS	84	A	12.640	3.478	-0.465
400	CE1	HIS	84	A	11.393	3.487	1.258
401	NE2	HIS	84	A	12.201	4.192	0.533
402	N	ALA	85	A	11.706	-1.690	0.180
403	CA	ALA	85	Α .	11.398	-2.240	1.463
404	С	ALA	85	A	12.553	-3.020	1.972
405	0	ALA	85	A	12.626	-3.221	3.161

Fig. 15.10

A1	om	Resi	duc	Chain	x	Y	Z
406	СВ	ALA	85	A	10.199	-3.130	1.402
407	N	ILE	86	A	13.538	-3.522	1.249
408	CA	ILE	86	A	14.606	-4.201	1.925
409	C	ILE	86	A	15.457	-3.071	2.472
410	0	ILE	86	A	15.835	-3.146	3.602
411	CB	ILE	86	A	15.300	-5.078 -6.058	0.903
412	CG1	ILE	86	A	14.276 16.470	-5.804	0.301 1.570
413	CG2	ILE	86	A	14.811	-7.212	-0.547
414	CD1	ILE	86 87	A A	15.835	-1.962	1.844
415	N	VAL	87	A	16.554	-0.892	2.511
416	CA	VAL	87	A	15.798	-0.536	3.762
417	C	VAL	87	A	16.284	-0.769	4.847
418	0	VAL VAL	87	A	16.672	0.325	1.574
419 420	CB CG1	VAL	87	A	16.890	1.606	2.333
421	CG2	VAL	87	A	17.858	0.133	0.660
422	N	GLN	88	A.	14.596	0.026	3.752
423	CA	GLN	88	A	13.942	0.411	4.993
424	C	GLN	88	A	13.900	-0.690	6.037
425	0	GLN	88	A	14.176	-0.400	7.186
426	CB	GLN	88	A	12.531	0.829	4.715
427	CG	GLN	88	A	12.099	2.208	5.146
428	CD	GLN	88	A	10.634	2.123	5.038
429	OE1	GLN	88	A	9.852	2.201	5.962
430	NE2	GLN	88	A	10.159	1.742	3.868
431	N	THR	89	A	13.582	-1.949	5.766
432	CA	THR	89	A	13.633	-2.920	6.799
433	С	THR	89	A	15.038	-3.073	7.309
434	ō	THR	89	A	15.197	-3.213	8.509
435	СВ	THR	89	A	13.156	-4.198	6.275
436	OG1	THR	89	A	11.863	-3.908	5.883
437	CG2	THR	89	A	12.973	-5.302	7.272
438	N	LEU	90	A	16.122	-3.061	6.519
439	CA	LEU	90	A	17.492	-3.003	7.017
440	С	LEU	90	A	17.782	-1.770	7.832
441	0	LEU	90	A	18.260	-1.850	8.947
442	CB	LEU	90	λ	18.454	-3.009	5.868
443	CG	LEU	90	A	19.904	-3.128	6.174
444	CD1	LEU	90	A	20.166	-4.389	6.967
445	CD2	LEU	90	A	20.641	-3.116	4.854
446	N	VAL	91	A	17.528	-0.569	7.354
447	CA	VAL	91	A	17.690	0.623	8.155
448	С	VAL	91	A	16.866	0.546	9.409
449	0	VAL	91	A	17.188	1.150	10.422
450	CB	VAL	91	A	17.282	1.803	7.334

Fig. 15.11

A	tom	Resi	due	Chain	x	Y	z
				,			
451	CG1	VAL	91	A	17.149	3.078	8.140
452	CG2	VAL	91	A	18.356	1.954	6.278
453	N	HIS	92	A	15.762	-0.193	9.453
454	CA	HIS	92	A	15.046	-0.309	10.719
455	С	HIS	92	λ	15.946	-0.915	11.723
456	0	HIS	92	λ	16.085	-0.444	12.816
457	СВ	HIS	92	A	13.805	-1.224	10.633
458	CG	HIS	92	A	12.902	-1.039	11.826
459	ND1	HIS	92	A	12.332	-1.848	12.711
460	CD2	HIS	92	A	12.540	0.216	12.159
461	CE1	HIS	92	A	11.678	-1.086	13.538
462	NE2	HIS	92	A	11.826	0.165	13.212
463	N	PHE	93	A	16.591	-1.984	11.374
464	CA	PHE	93	A	17.374	-2.777	12.262
465	C	PHE	93	A	18,620	-2.032	12.652
466	0	PHE	93	A	19.005	-1.893	13.796
467	CB	PHE	93	A	17.592	-4.024	11.472
468	CG	PHE	93	A	18.741	-4.858	11.921
469	CD1	PHE	93	A	20.039	-4.461	11.662
470	CD2	PHE	93	A	18.474	-6.111	12.455
471	CE1	PHE	93	A	21.058	-5.341	11.903
472	CE2	PHE	93	A	19.507	-6.993	12.676
473	CZ	PHE	93	A	20.790	-6.600	12.394
474	N	ILE	94	A	19.399	-1.476	11.751
475	CA	ILE	94	A	20.477	-0.592	12.182
476	C	ILE	94	A	19.938	0.454	13.131
477	0	ILE	94	A	20.706	1.117	13.787
478	CB	ILE	94	A	21.036	-0.005	10.907
479	CG1	ILE	94	A	21.808	-1.156	10.367
480	CG2	ILE	94	A	21.876	1.246	11.008
481	CD1	ILE	94	A	22.103	-0.863	8.903
482	N	ASN	95	A	18.635	0.704	13.302
483	CA	asn	95	A	18.237	1.955	13.901
484	С	ASN	95	A	16.753	2.185	14.183
485	0	asn	95	A	16.082	3.112	13.702
486	CB	ASN	95	A	18.719	3.074	13.024
487	CG	ASN	95	A	18.687	4.221	13.977
488	OD1	ASN	95	A	18.243	4.291	15.040
489	ND2	ASN	95	A	19.771	5.172	13.602
490	N	PRO	96	A	16.204	1.357	15.006
491	CA	PRO	96	A	14.787	1.108	15.108
492	C	PRO	96	A	13.919	2.281	15.324
493	0	PRO	96	A	12.713	2.226	15.089
494	CB	PRO	96	λ	14.647	0.084	16.217
495	CG	PRO	96	A	16.039	-0.490	16.300

Fig. 15.12

A	tom	Re	siduc	Chain	x	Y	<b>z</b>
496	CD	PRO	96	A	16.846	0.801	16.173
497	N	GLU		A	14.559	3.336	15.774
498	CA	GLU	97	A	13.846	4.558	15.992
499	С	GLU		A	14.019	5.333	14.722
500	0	GLU	97	A	13.031	5.781	14.197
501	CB	Gra	97	A	14.429	5.329	17.184
502	CG	GLU	97	A	13.550	5.172	18.453
503	CD	GLU	97	A	13.769	3.807	19.163
504	OE1	GLU	97	A	14.723	3.749	19.999
505	OE2	GLU	97	A	12.998	2.824	18.896
506	И	THR	98	A	15.145	5.584	14.080
507 508	CA	THR	98	A	15.073	6.445	12.897
509	C O	THR THR	98 98	A A	14.148	6.043	11.715
510	CB	THR	98	A	14.015	6.906	10.851
511	OG1	THR	98	A	16.514 17.067	6.624	12.388
512	CG2	THR	98	A	17.451	5.331 7.417	12.373 13.322
513	N	VAL	99	A	13.482	4.877	11.518
514	CA	VAL	99	A	12.654	4.635	10.334
515	C	VAL	99	A	11.675	3.508	10.660
516	o	VAL	99	A	12.183	2.611	11.310
517	CB	VAL	99	A	13.546	4.251	9.139
518	CG1	VAL	99	A	13.636	2.753	8.774
519	CG2	VAL	99	A	12.902	4.955	7.978
520	N	PRO	100	A	10.408	3.461	10.322
521	CA	PRO	100	A	9.442	2.427	10.687
522	С	PRO	100	A	9.639	1.132	10.021
523	0	PRO	100	A	10.611	1.033	9.342
524	CB	PRO	100	A	8.119	3.003	10.359
525	CG	PRO	100	A	8.348	4.485	10.464
526	CD		100	A	9.713	4.645	9.854
527	N		101	A	8.889	0.042	10.071
528	CA	LYS		A	9.245	-1.083	9.226
529	C	LYS		A	8.344	-0.893	8.022
530	0	LYS		A	7.361	-0.183	8.151
531	CB	LYS		A	8.901	-2.426	9.859
532	CG	LYS		A	8.737	-2.477	11.364
533	CD	LYS		A	8.039	-3.748	11.889
534 535	CE	LYS		A	6.454	-3.645	12.044
536	NZ	LYS		Α	5.616	-3.935	10.845
537	N CA	PRO PRO		A	8.452	-1.381	6.846
538	CA	PRO		A A	7.556 6.203	-1.032 -1.549	5.775
539	0	PRO		λ	6.087	-2.464	6.093 6.892
540	CB	PRO		Ä	8.171	-1.646	4.535

Fig. 15.13

A	tom	Residue	Chain	X	<u>Y</u>	Z
541	CG	PRO 102	A	8.812	-2.855	5.077
542	CD	PRO 102	A	9.333	-2.458	6.451
543	N	CYS 103	A	5.086	-1.078	5.549
544	CA	CYS 103	A	3.887	-1.780	5.921
545	С	CYS 103	A	3.229	-2.407	4.753
546	0	CYS 103	A	3.592	-2.289	3.598
547	CB	CYS 103	· <b>A</b>	2.930	-0.840	6.599
548	SG	CYS 103	A	2.131	-0.050	5.233
549	N	CYS 104	A	2.191	-3.102	5.117
550	CA	CYS 104	A	1.736	-4.199	4.338
551	C	CYS 104	A	0.466	-3.786	3.641
552	0	CYS 104	A	-0.555	-3.520	4.240
553	CB	CYS 104	A	1.633	-5.221	5.372
554	SG	CYS 104	A	0.823	-6.586	4.674
555	N	ALA 105	A	0.490	-3.719	2.337
556	CA	ALA 105	A	-0.396	-2.886	1.605
557	С	ALA 105	A	-0.839	-3.742	0.482
558	0	ALA 105	A	-0.170	-4.745	0.282
559	CB	ALA 105	A	0.400	-1.708	1.149
560	N	PRO 106	A	-1.857	-3.510	-0.271
561	CA	PRO 106	A	-2.156	-4.293	-1.435
562	С	PRO 106	A	-1.352	-3.905	-2.675
563	0	PRO 106	A	-1.065	-2.765	-2.987
564	CB	PRO 106	A	-3.647	-4.136	-1.574
565	CG	PRO 106	A	-3.879	-2.759	-1.095
566	CD	PRO 106	<b>A</b> .	-3.107	-2.927	0.186
567	N	THR 107	Α	-0.999	-4.955	-3.391
568	CA	THR 107	A	-0.073	-4.974	-4.479
569	С	THR 107	Α	-1.001	-4.913	-5.659
570	0	THR 107	A	-0.955	-3.955	-6.424
571	СВ	THR 107	A	0.666	-6.265	-4.172
572	OG1	THR 107	A	1.728	-5.736	-3.391
573	CG2	THR 107	A	1.115	-7.153	-5.311
574	N	GLN 108	λ	-1.900	-5.847	-5.940
575	CA	GLN 108	A	-2.913	-5.442	-6.865
576	С	GLN 108	A	-4.352	-5.682	-6.500
577	0	GLN 108	A	-4.739	-6.674	-5.905
578	CB	GLN 108	A	-2.460	-6.080	-8.189
579	CG	GLN 108	A	-2.948	-7.440	-8.663
580	CD	GLN 108	A	-2.234	-8.459	-7.886
581	OE1	GLN 108	A	-1.660	-8.222	-6.832
582	NE2	GLN 108	A	-2.298	-9.656	-8.471
583	N	LEU 109	A	-5.100	-4.662	-6.917
584	CA	LEU 109	A	-6.522	-4.430	-6.644
585	С	LEU 109	A	-7.495	-4.780	-7.798

Fig. 15.14

A	tom	Residue	Chain	<u>x</u>	Y	
			_			
586	0	LEU 109	A	-7.256	-4.573	-8.986
587	CB	LEU 109	A	-6.694	-2.932	-6.225
588	CG	LEU 109	A	-6.102	-2.671	-4.830
589	CD1	LEU 109	A	-6.012	-1.216	-4.502
590	CD2	LEU 109	A	-7.022	-3.304	-3.811
591	N	ASN 110	A	-8.659	-5.343	-7.521
592	CA	ASN 110	A	-9.570	-5.629	-8.584
593	C	ASN 110	A	-10.824	-4.870	-8.418
594	0	ASN 110	A	-10.985	-4.117	-7.471
595	CB	ASN 110	A	-9.945	-7.054	-8.620
596	CG	ASN 110 ASN 110	A A	-8.771	~7.863	-9.017
597	OD1 ND2	ASN 110	A	-8.810 -7.697	-9.080	-8.781
598 500	ND2 N	ALA 111	A	-7.687 -11.767	-7.281 -5.052	-9.560
599 600	CA	ALA 111	A	-12.876	-4.131	-9.338
601	C	ALA 111	A	-14.026	-4.131	-9.476 -9.121
602	0	ALA 111	A	-13.880	-6.178	-9.121 -9.05B
603	СВ	ALA 111	A	-12.993		-10.928
604	N	ILE 112	A	-15.182	-4.401	-8.876
605	CA	ILE 112	A	-16.371	-5.172	-8.539
606	C	ILE 112	A	-17.371	-4.462	-9.404
607	0	ILE 112	A	-17.265	-3.251	-9.506
60B	CB	ILE 112	A	-16.583	-5.062	-6.980
609	CG1	ILE 112	A	-16.477	-6.535	-6.632
610	CG2	ILE 112	A	-17.869	-4.480	-6.379
611	CD1	ILE 112	A	-17.762	-7.334	-6.966
612	N	SER 113	A	-18.320		-10.037
613	CA	SER 113	A	-19.389		-10.673
614	С	SER 113	A	-20.670	-4.529	-9.920
615	0	SER 113	A	-20.965	-5.655	-9.564
616	СВ	SER 113	A	-19.576		-12.074
617	OG	SER 113	A	-18.826	-4.052	-12.946
618	N	VAL 114	A	-21.472	-3.484	-9.631
619	CA	VAL 114	A	-22.751	-3.722	-8.970
620	С	VAL 114	A	-23.830	-3.148	-9.882
621	0	VAL 114	A	-23.658	-2.119	-10.522
622	CB	VAL 114	A	-22.897	-3.057	-7.536
623	CG1	VAL 114	A	-21.637	-3.226	-6.742
624	CG2	VAL 114	A	-23.194	-1.588	-7.611
625	N	LEU 115	A	-24.960	-3.847	-9.955
626	CA	LEU 115	A	-26.160	-3.492	-10.709
627	С	LEU 115	A	-27.151	-2.699	-9.903
628	0	LEU 115	A	-27.603	-3.173	-8.860
629	СВ	LEU 115	A	~26.840	-4.789	-11.194
630	CG	LEU 115	A	-28.187	-4.790	-11.902

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Fig. 15.15

A	tom	Residue	Chain	X	Y	Z
631	CD1	LEU 115	A	-28.011	-4.403	-13.339
632	CD2	LEU 115	A	-28.797		-11.806
633	N	TYR 116	A	-27.598		-10.254
634	CA	TYR 116	A	-28.495	-0.859	-9.329
635	C	TYR 116	λ	-29.472	0.124	-9.908
636	0	TYR 116	A	-29.241		-11.014
637 638	CB CG	TYR 116	A	-27.619		-8.316
639	CD1	TYR 116 TYR 116	A A	-26.890	0.935	-8.941
640	CD2	TYR 116	A	-25.664 -27.474	0.712 2.166	-9.513
641	CE1	TYR 116	A	-25.034	1.750	-8.888 -10.131
642	CE2	TYR 116	A	-26.850	3.205	-9.502
643	cz	TYR 116	A	-25.669		-10.141
644	OH	TYR 116	A	-25.140		-10.902
645	N	PHE 117	A	-30.566	0.506	-9.242
646	CA	PHE 117	A	-31.478	1.445	-9.867
647	С	PHE 117	A	-30.914	2.785	-9.556
648	0	PHE 117	A	-30.775	3.080	-8.381
649	CB	PHE 117	A	-32.882	1.514	-9.284
650	CG	PHE 117	A	-33.447	0.159	-8.982
651	CD1	PHE 117	A	-33.259	-0.395	-7.738
652	CD2	PHE 117	A	-34.137	-0.534	-9.952
653	CE1	PHE 117	A	-33.750	-1.660	-7.482
654	CE2	PHE 117	A	-34.618	-1.807	-9.709
655	CZ	PHE 117	A	-34.419	-2.373	-8.466
656	N	ASP 118	A	-30.555		-10.458
657	CA	ASP 118	A	-30.375	5.050	-9.964
658	С	ASP 118	A	-31.794	5.566	-9.803
659	0	ASP 118	A	-32.746	4.804	-9.876
660	CB	ASP 118	A	-29.612		-10.973
661 662	CG	ASP 118	A	-30.317		-12.316
663	OD1 OD2	ASP 118 ASP 118	A A	-29.855		-13.211
664	N	ASP 118	A	-31.322 -31.939	6.868	-12.467 -9.594
665	CA	ASP 119	A	-31.939	7.509	-9.594 -9.718
666	C	ASP 119	A	-33.216		-11.070
667	0	ASP 119	A.	-33.359		-11.946
668	СВ	ASP 119	A	-33.033	9.030	-9.618
669	CG	ASP 119	A.	-31.942		-10.598
670	OD1	ASP 119	A	-31.911		-11.719
671	OD2	ASP 119	A	-31.115		-10.232
672	N	SER 120	A	-35.161		-11.218
673	CA	SER 120	A	-36.125		-12.146
674	С	SER 120	A	-36.295		-11.621
675	0	SER 120	A	-36.515	5.484	-10.431

Fig. 15.16

Atom	Residuc	Chain	x	Y	Z
676 CB	SER 120	A	- 25 500	2 057	10 504
677 OG	SER 120	A	-35.506		-13.534
678 N	SER 121		-35.100		-14.079
679 CA	SER 121	A A	-36.217		-12.388
680 C	SER 121	A	-35.537		-11.785
681 0	SER 121	Ä	-34.705 -34.958		-12.949 -13.464
682 CB	SER 121	A	-36.551		-13.464
683 OG	SER 121	A	-37.674		-10.810
684 N	ASN 122	A	-33.701		-13.473
685 CA	ASN 122	A	-32.990		-14.519
686 C	ASN 122	A	-32.307		-13.858
687 0	ASN 122	A	-31.660		-12.857
688 CB	ASN 122	A	-31.913		-15.157
689 CG	ASN 122	A	-32.456		-16.232
690 OD1	ASN 122	A A	-33.543		-16.193
691 ND2	ASN 122	A	-31.577		-17.223
692 N	VAL 123	A	-32.362		-14.271
693 CA	VAL 123	A	-31.447	-0.312	
694 C	VAL 123	A	-30.061	-0.155	
695 O	VAL 123	A	-29.858	-0.450	
696 CB	VAL 123	A	-31.992	-1.647	
697 CG1	VAL 123	A	-31.033	-2.769	
698 CG2	VAL 123	A	-33.230	-1.785	
699 ห	ILE 124	A	-29.010		-13.637
700 CA	ILE 124	A	-27.704		-14.288
701 C	ILE 124	A	-26.583	-0.682	-13.814
702 0	ILE 124	A	-26.689	-1.441	-12.866
703 CB	ILE 124	A	-27.187	1.630	-14.333
704 CG1	ILE 124	A	-27.086	2.273	-13.025
705 CG2	ILE 124	A	-28.198	2.409	-15.116
706 CD1	ILE 124	A	-26.681	3.705	-13.362
707 N	LEU 125	A	-25.411	-0.711	-14.431
708 CA	LEU 125	Ä	-24.372	-1.639	-13.989
709 C	LEU 125	A	-23.210	-0.681	-13.840
710 0	LEU 125	A	-22.730	-0.263	-14.888
711 CB	LEU 125	A	-24.075	-2.659	-15.069
712 CG	LEU 125	A	-23.261	~3.900	-14.761
713 CD1	LEU 125	A	-23.927	-4.875	-13.791
714 CD2	LEU 125	A	-23.108	-4.618	-16.093
715 N	LYS 126	A	-22.754	-0.305 ·	-12.615
716 CA	LYS 126	A	-21.529	0.463	-12.487
717 C	LYS 126	A	-20.416	-0.467	-12.106
718 O	LYS 126	A	-20.595	-1.572 -	-11.608
719 CB	LYS 126	A	-21.624	1.556 -	-11.437
720 CG	LYS 126	A	-20.629	2.603 -	-12.021

Fig. 15.17

At	om	Residuc	Chain	<u> </u>	<u> Y</u>	Z
201	CD	LYS 126	A	-20.367	3.908	-11.203
721 722	CD CE	LYS 126	A	-19.079		-11.635
723	NZ	LYS 126	A	-17.843		-11.529
724	N	LYS 127	λ	-19.199		-12.352
725	CA	LYS 127	A	-18.017		-12.143
726	C	LYS 127	A	-17.282	0.026	-11.185
727	0	LYS 127	A	-17.184	1.224	-11.370
728	CB	LYS 127	A	-17.097	-0.968	-13.339
729	CG	LYS 127	A	-16.235	-2.196	-13.072
730	CD	LYS 127	Α	-15.334	-2.558	-14.231
731	CE	LYS 127	A	-14.497	-1.365	-14.645
732	NZ	LYS 127	А	-13.360	-1.909	~15.350
733	N	TYR 128	A	-16.763	-0.565	-10.145
734	CA	TYR 128	A	-16.120	0.206	-9.143
735	С	TYR 128	А	-14.784	-0.346	-9.316
736	0	TYR 128	A	-14.664	-1.559	-9.483
737	CB	TYR 128	A	-16.692	-0.131	-7.789
738	CG	TYR 128	A	-17.935	0.643	-7. <b>63</b> 6
739	CD1	TYR 128	A	-19.083	0.008	-7.229
740	CD2	TYR 128	A	-17.936	1.966	-7.982
741	CE1	TYR 128	A	-20.272	0.703	-7.216
742	CE2	TYR 128	A	-19.101	2.672	-7.977
743	CZ	TYR 128	A	-20.270	2.038	-7.613
744	OH	TYR 128	A	-21.474	2.754	-7.674
745	N	ARG 129	A	-13.741	0.463	-9.292
746	CA	ARG 129	A	-12.483	-0.169	-9.488
747	С	ARG 129	A	-11.895	-0.158	-8.145
748	0	ARG 129	A	-12.287	0.615	-7.288
749	CB	ARG 129	A	-11.616	0.626	
750	CG	ARG 129	A	-12.191		-11.208
751	CD	ARG 129	A	-11.134	2.214	
752	NE	ARG 129	A	-10.607	1.001	
753	CZ	ARG 129	A	-11.262	0.401	
754	NH1	ARG 129	A	-12.339		-14.529
755	NH2	ARG 129	A	-10.849	-0.B46	
756	N	ASN 130	A	-10.935	-1.039	-7.949
757	CA	ASN 130	A	-10.085	-1.012	-6.790
758	С	ASN 130	A	-10.851	-1.277	-5.599
759	0	ASN 130	A	-10.708	-0.669	-4.572
760	CB	ASN 130	A	-9.393	0.323	-6.609
761	CG	ASN 130	A	-8.261	0.341	-7.600
762	OD1	ASN 130	A	-7.739	-0.694	-7.981
763	ND2	ASN 130	A	-7.853	1.492	-8.122
764	N	MET 131	A	-11.728	-2.248	-5.704
765	CA	MET 131	A	-12.531	-2.586	-4.577

Fig. 15.18

At	om	Residue	Chain	<u> </u>	<u>Y</u>	
766	c	MET 131	Α	-12.155	-3.895	-3.926
767	o	MET 131	A	-12.642	-4.228	-2.858
768	СВ	MET 131	A	-13.944	-2.576	-5.083
769	CG	MET 131	A ·	-14.347	-1.149	-5.256
770	SD	MET 131	A	-15.527	-0.836	-3.944
771	CE	MET 131	A	-14.884	0.789	-3.671
772	N	VAL 132	A	-11.291	-4.690	-4.521
773	CA	VAL 132	A	-11.025	-6.042	-4.048
774	C	VAL 132	A	-9.531	-6.189	-3.925
775	o	VAL 132	A	-8.827	-5.797	-4.819
776	СВ	VAL 132	A	-11.556	-7.015	-5.070
777	CG1	VAL 132	A	-11.478	-8.446	-4.622
778	CG2	VAL 132	A	-13.002	-6.636	-5.321
779	N	VAL 133	A	-8.890	-6.710	-2.915
780	CA	VAL 133	A	-7.469	-6.920	-2.922
781	С	VAL 133	A	-7.287	-8.237	-3.624
782	0	VAL 133	A	-8.076	-9.130	-3.369
783	CB	VAL 133	A	-7.014	-6.965	-1.472
784	CG1	VAL 133	λ	-5.786	-7.825	-1.195
785	CG2	VAL 133	A	-6.799	-5.522	-1.123
786	N	ARG 134	A	-6.278	-8.372	-4.493
787	CA	ARG 134	A	-5.826	-9.648	-5.013
788	С	ARG 134	A	-4.558	-10.087	-4.286
789	0	ARG 134	Α	-4.366	-11.235	-3.880
790	CB	ARG 134	A	-5.517	-9.521	-6.505
791	CG	ARG 134	A	-6.403	-10.251	-7.540
792	CD	ARG 134	A	-5.677	-10.152	-8.920
793	NE	ARG 134	A	-6.330	-10.776	-10.106
794	CZ	ARG 134	A	-6.819	-12.047	-10.211
795	NH1	ARG 134	A	-7.257	-12.490	-11.438
796	NH2	ARG 134	A	-6.948	-12.890	-9.134
797	N	ALA 135	A	-3.591	-9.198	~4.086
798	CA	ALA 135	A	-2.487	-9.628	-3.277
799	c	ALA 135	A	-1.918	-8.469	-2.523
800	0	ALA 135	A	-2.177	-7.321	-2.854
801	CB	ALA 135	A	-1.466	-10.268	-4.196
802	n	CYS 136	A	-1.126	-8.787	-1.485
803	CA	CYS 136	A	-0.495	-7.839	-0.552
804	C	CYS 136	A	1.009	-8.034	-0.432
805	ō	CYS 136	A	1.465	-9.160	-0.589
806	CB	CYS 136	λ	-0.883	-7.996	0.872
807	SG	CYS 136	A	-2.530	-8.641	1.107
808	N	GLY 137	A	1.790	-7.010	-0.149
809	CA	GLY 137	A	3.125	-7.303	0.230
	C	GLY 137	A	3.804	-6.043	0.544

Fig. 15.19

A	lom	Res	idue	Chain	x	Y	<u>z</u>
811	0	GLY		A	3.166	-5.012	0.413
812	N		138	A	5.077	-6.032	0.947
813	CA		138	A	5.617	-4.923	1.740
814	С		138	A	6.031	-3.785	0.894
815	0		138	A	6.745	-4.136	-0.019
816	CB		138	A	6.815	-5.368	2.504
817	SG		138	A	6.442	-7.019	3.133
818	N		139	A	5.732	-2.511	1.016
819	CA		139	A	6.451	-1.586	0.176
820	С	HIS	139	A	6.716	-0.472	1.145
821	0		139	A	5.993	-0.333	2.154
822	CB		139	A	5.651	-1.066	-1.031
823	CG		139	A	5.126	-2.180	-1.889
824	ND1		139	A	4.592	-3.381	-1.428
825	CD2	HIS	139	A	5.565	-2.289	-3.201
826	CE1	HIS	139	A	4.749	-4.140	-2.529
827	NE2	HIS	139	A	5.322	-3.570	-3.624
828	OXT	HIS	139	A	7.321	0.362	0.777
829	N	GLN	36	В	-1.316	13.234	11.930
830	CA	GLN	36	В	-0.019	13.097	11.315
831	С	GLN	36	B	-0.372	12.092	10.267
832	0	GLN	36	В	-0.507	10.903	10.466
833	CB	GLN	36	В	0.994	12.511	12.324
834	CG	GLN	36	В	0.822	10.996	12.674
835	CD	GLN	36	В	-0.460	10.531	13.376
836	OE 1	GLN	36	В	-1.426	11.270	13.478
837	NE2	GLN	36	В	-0.520	9.334	13.901
838	N	ALA	37	В	-0.523	12.730	9.128
839	CA	ALA	37	В	-1.432	12.344	8.073
840	С	ALA	37	В	-0.603	11.520	7.202
841	0	ALA	37	В	0.573	11.779	7.257
842	CB	ALA	37	В	-1.894	13.588	7.326
843	N	CYS	38	В	-0.971	10.577	6.394
844	CA	CYS	38	В	0.057	9.856	5.706
845	С	CYS	38	В	0.697	10.748	4.703
846	0	CYS	38	В	-0.011	11.155	3.781
847	CB	CYS	38	B	-0.585	8.677	5.078
848	SG	CYS	38	B	0.292	8.306	3.618
849	N	LYS	39	В	2.000	11.027	4.892
850	CA	LYS	39	В	2.823	11.772	3.937
851	С	LYS	39	В	4.228	11.167	3.694
852	0	LYS	39	В	4.679	10.270	4.389
853	CB	LYS	39	В	2.990	13.236	4.403
854	CG	LYS	39	В	3.093	13.354	5.921
855	CD	LYS	39	В	3.434	14.707	6.546

Fig. 15.20

A	tom	Resi	due	Chain	x	Y	Z
856	CE	LYS	39	В	3.222	14 463	0.000
857	NZ	LYS	39	В	3.429	14.463	8.066
858	N	LYS	40	В	4.966	15.675 11.640	8.873
859	CA	LYS	40	В	6.314	11.192	2.693 2.434
860	C	LYS	40	В	7.224	11.807	3.433
861	o	LYS	40	В	6.889	12.880	3.892
862	CB	LYS	40	В	6.814	11.640	1.096
863	CG	LYS	40	В	8.068	10.922	0.629
864	CD	LYS	40	В	8.867	11.865	-0.243
865	CE	LYS	40	В	8.006	12.488	-1.310
866	NZ	LYS	40	В	8.797	13.441	-2.059
867	N	HIS	41	В	8.361	11.220	3.821
868	CA	HIS	41	В	9.283	11.844	4.760
869	С	HIS	41	В	10.701	11.549	4.352
870	0	HIS	41	В	10.914	10.791	3.425
871	CB	HIS	41	В	9.179	11.290	6.141
872	CG	HIS	41	В	7.860	11.488	6.797
873	ND1	HIS	41	В	7.584	12.100	7.948
874	CD2	HIS	41	В	6.778	10.752	6.444
875	CE1	HIS	41	В	6.404	11.711	8.330
876	NE2	HIS	41	В	5.937	10.898	7.415
877	N	GLU	42	В	11.751	12.069	4.966
878	CA	GLU	42	В	13.094	11.965	4.415
879	С	GLU	42	B	13.878	10.810	4.901
880	0	GLU	42	В	13.613	10.301	5.969
881	CB	GLU	42	В	13.875	13.207	4.718
882	ÇG	GLU	42	В	13.369	14.277	3.745
883	CD	GLU	42	В	14.490	15.264	3.400
884	OE1	GLU	42	В	14.161	16.446	3.123
885	OE2	GLU	42	В	15.681	14.846	3.405
886	N	LEU	43	В	14.868	10.303	4.208
887	CA	LEU	43	В	15.662	9.220	4.794
888	С	LEU	43	В	16.791	9.303	3.810
889	0	LEU	43	В	16.590	9.000	2.644
890	CB	LEU	43	В	15.082	7.834	4.628
891	CG	LEU	43	В	15.121	6.808	5.726
892	CD1	LEU	43	В	15.317	5.447	5.064
893	CD2	LEU	43	В	16.201	7.098	6.734
894	N	TYR	44	В	17.960	9.706	4.244
895	CA	TYR	44	В	19.116	9.691	3.414
896	С	TYR	44	В	19.731	8.374	3.684
897	0	TYR	44	В	20.074	8.206	4.838
898	CB	TYR	44	В	19.987	10.792	3.853
899	CG	TYR	44	В	21.075	10.941	2.875
900	CD1	TYR	44	В	22.293	10.457	3.244

Fig. 15.21

At	om	Resid	uc	Chain	X	Y	<u>z</u>
			44	В	20.850	11.476	1.648
901	CD2	TYR	44	В	23.318	10.366	2.365
902	CE1	TYR	44	В	21.874	11.387	0.757
903	CE2	TYR TYR	44	В	23.066	10.774	1.099
904	CZ	TYR	44	B	24.015	10.452	0.125
905	OH	VAL	45	В	19.912	7.407	2.784
906	N CA	VAL	45	В	20.665	6.229	3.164
907	C	VAL	45	В	22.087	6.437	2.654
908 909	0	VAL	45	В	22.279	6.805	1.507
910	CB	VAL	45	В	20.035	5.032	2.545
911	CG1	VAL	45	В	20.695	3.749	2.995
912	CG2	VAL	45	В	18.586	5.039	2.976
913	N	SER	46	В	23.114	6.225	3.477
914	CA	SER	46	В	24.493	6.439	3.107
915	c	SER	46	В	25.038	5.102	2.862
916	0	SER	46	В	24.941	4.287	3.750
917	CB	SER	46	В	25.291	7.048	4.229
918	OG	SER	46	В	26.655	7.224	3.852
919	N	PHE	47	В	25.629	4.746	1.749
920	CA	PHE	47	В	25.991	3.376	1.514
921	С	PHE	47	В	26.978	2.852	2.491
922	0	PHE	47	В	27.217	1.663	2.537
923	CB	PHE	47	В	26.594	3.221	0.184
924	CG	PHE	47	В	25.580	3.521	-0.868
925	CD1	PHE	47	В	24.477	2.747	-0.984
926	CD2	PHE	47	В	25.878	4.449	-1.823
927	CE1	PHE	47	В	23.731	2.826	-2.115
928	CE2	PHE	47	B	25.135	4.505	-2.974
. 929	CZ	PHE	47	В	24.071	3.673	-3.132
930	N	ARG	48	В	27.608	3.672	3.320
931	CA	ARG	48	В	28.339	3.167	4.458
932	С	ARG	48	В	27.398	2.413	5.320
933	Ο,	ARG	48	В	27.646	1.284	5.611
934	CB	ARG	48	В	28.954	4.292	5.278 4.494
935	CG	ARG	48	В	30.046	5.035	5.396
936	CD	ARG	48	В	30.896	5.928	4.756
937	NE	ARG	48	В	32.154	6.274	4.756
938	CZ	ARG	48	В	32.651	7.514	5.519
939	NH1	ARG	48	В	31.950	8.482 7.849	4.304
940	NH2	ARG	48	B -	33.859	2.920	5.771
941	N	ASP	49	В	26.272 25.514	2.260	6.816
942	CA	ASP	49	В	24.987	0.889	6.405
943	С	ASP	49	В	24.545	0.069	7.199
944	0	ASP	49	В	•	3.105	7.229
945	CB	ASP	49	В	24.328	3.105	1.223

Fig. 15.22

At	om	Resid	luc	Chain	x	Y	Z
		_		_	24.718	4.526	7.549
946	CG	ASP	49	B	23.837	5.362	7.347
947	OD1	ASP	49	В	25.851	4.840	7.969
948	OD2	ASP	49	В	25.005	0.563	5.125
949	N	LEU	50	В	24.555	-0.751	4.716
950	CA	LEU	50	В	25.746	-1.589	4.430
951	С	LEU	50	В	25.685	-2.699	3.895
952	0	LEU	50	B	23.706	-0.649	3.461
953	CB	LEU	50	В	22.524	0.294	3.611
954	CG	LEU	50	В	22.301	0.924	2.257
955	CD1	LEU	50	B B	21.289	-0.430	4.147
956	CD2	LEU	50	В	26.887	-1.028	4.780
957	N	GLY	51	B	28.143	-1.664	4.485
958	CA	GLY	51	. 1B	28.483	-1.605	3.017
959	C	GLY	51	. B	29.593	-1.964	2.642
960	0	GLY	51 52	B	27,628	-1.178	2.086
961	N	TRP	52	В	28.069	-1.113	0.692
962	CA	TRP	52	В	29,287	-0.215	0.418
963	C	TRP	52	В	29.372	0.385	-0.629
964	0	TRP	52 52	В	26.856	-0.656	-0.156
965	CB	TRP TRP	52	В	25.726	-1.702	-0.123
966	CG CD1	TRP	52	В	25.857	-2.997	0.341
967 968	CD2	TRP	52	В	24.335	-1.394	-0.582
969	NE1	TRP	52	В	24.672	-3.554	0.236
970	CE2	TRP	52	В	23.731	-2.740	-0.270
971	CE3	TRP	52	В	23.505	-0.402	-1.076
972	CZ2	TRP	52	В	22.376	-2.951	-0.421
973	CZ3	TRP	52	В	22.157	-0.672	-1.220
974	CH2	TRP	52	В	21.604	-1.908	-0.885
975	N .	GLN	53	В	30.313	-0.015	1.236
976	CA	GLN	53	18	30.998	1.273	1.141
977	C	GIN	53	В	32.040	1.358	0.116
978	o	GLN	53	В	32.702	2.374	-0.014
979	СВ	GLN	53	В	31.742	1.728	2.405
980	CG	GLN	53	В	31.993	0.649	3.446
981	CD	GLN	53	В	33.274	0.967	4.237
982	OE1	GLN	53	В	34.189	0.121	4.274
983	NE2	GLN	53	В	33.399	2.156	4.889
984	N	ASP	54	В	32.235	0.306	-0.647
985	CA	ASP	54	В	33.600	-0.101	-0.948
986	C	ASP	54	В	33.381	-0.824	-2.214
987	o	ASP	54	В	33.245	-2.036	-2.082
988		ASP	54	В	34.116	-1.083	0.114
989		ASP	54	В	32.963	-1.909	0.747
990		ASP	54	В	33.101	-2.317	1.928

Fig. 15.23

A	tom	Resi	due	Chain	x	Y	z
991	OD2	ASP	54	В	31.940	-2.112	0.062
992	N	TRP	55	В	33.314	-0.266	-3.427
993	CA	TRP	55	В	32.570	-1.004	-4.476
994	С	TRP	55	В	32.019	-0.003	-5.421
995	0	TRP	55	В	32.165	-0.187	-6.621
996	CB	TRP	55	В	31.258	-1.689	-4.123
997	CG	TRP	55	В	31.339	-3.126	-3.756
998	CD1	TRP	55	В	32.336	-4.015	-4.070
999	CD2	TRP	55	В	30.132	-3.715	-3.105
1000	NE1	TRP	55	В	31.855	-5.199	-3.695
1001	CE2	TRP	55	В	30.594	-5.172	-3.148
1002	CE3	TRP	55	В	28.951	-3.382	-2.460
1003	CZ2	TRP	55	В	29.753	-6.141	-2.581
1004	CZ3	TRP	55	В	28.177	-4.393	-1.896
1005	CH2	TRP	55	В	28.549	-5.738	-1.967
1006	N	ILE	56	B	31.399	1.020	-4.769
1007	CA	ILE	56	В	30.572	2.056	-5.356
1008	С	ILE	56	В	31.459	3.266	-5.650
1009	0	ILE	56	B	32.148	3.707	-4.741
1010	CB	ILE	56	B	29.469	2.363	-4.346
1011	CG1	ILE	56	B	28.247	1.655	-4.774
1012	CG2	ILE	56	В	29.164	3.835	-4.244
1013	CD1	ILE	56	В	28.212	0.333	-4.085
1014	N	ILE	57	B B	31.511 32.213	3.841 5.100	-6.85 <b>7</b> -7.039
1015	CA	ILE	57 57		31.178	6.206	-6.753
1016	C	ILE		В	31.414	7.392	-6.654
1017	0	ILE	57 57	B B	32.735	5.162	-8.493
1018	CB	ILE		В	33.659		
1019	CG1	ILE	57 57	В	33.527	4.001 6.433	-8.726 -8.784
1020	CG2	ILE	57		34.001		-10.220
1021	CD1	ILE	58	B B	29.934	5.831	-6.570
1022	N	ALA	58	В	28.802	5.950	-7.504
1023 1024	CA C	ALA ALA	58	. B	28.040	7.166	-7.145
1025	0	ALA	58	B	28.524	8.150	-7.690
1025	СВ	ALA	58	В	27.934	4.732	-7.338
1027	N	PRO	59	В	26.999	7.188	-6.358
1028	CA	PRO	59	В	26.820	8.263	-5.382
1029	C	PRO	59	В	27.333	7.862	-4.008
1030	o	PRO	59	В	27.512	6.69B	-3.744
1031	СВ	PRO	59	В	25.317	8.611	-5.384
1032	CG	PRO	59	В	24.697	7.280	-5.680
1033	CD	PRO	59	В	25.659	6.734	-6.707
1034	N	GLU	60	В	27.622	8.703	-3.030
1035	CA	GLU	60	В	27.941	8.146	-1.719

Fig. 15.24

1036 C GLU 60 B 26.735 7.734 -0.926 1037 O GLU 60 B 26.813 7.132 -0.867 1039 CG GLU 60 B 28.745 9.151 -0.867 1039 CG GLU 60 B 27.995 10.280 -0.117 1040 CD GLU 60 B 27.231 9.868 1.144 1041 OE1 GLU 60 B 27.559 8.834 1.775 1042 OE2 GLU 60 B 26.329 10.634 1.518 1043 N GLY 61 B 25.552 8.049 -1.395 1044 CA GLY 61 B 23.253 8.252 -1.519 1046 C GLY 61 B 23.253 8.252 -1.519 1046 C GLY 61 B 23.535 8.769 -2.595 1047 N TYR 62 B 22.021 8.246 -1.038 1048 CA TYR 62 B 20.934 8.777 -1.830 1049 C TYR 62 B 19.767 9.046 -0.932 1050 O TYR 62 B 19.767 9.046 -0.932 1050 C TYR 62 B 19.767 9.046 -0.932 1050 C TYR 62 B 19.833 6.572 -2.370 1053 CD1 TYR 62 B 20.425 5.395 -1.917 1055 CE1 TYR 62 B 19.624 4.316 -1.563 1056 CE2 TYR 62 B 19.624 4.316 -1.563 1057 CZ TYR 62 B 19.624 4.316 -1.563 1058 CH TYR 62 B 19.624 4.316 -1.563 1059 N ALA 63 B 18.720 9.711 -1.407 1060 CA ALA 63 B 18.720 9.711 -1.407 1060 CA ALA 63 B 16.485 8.722 -0.918 1062 O ALA 63 B 16.485 8.722 -0.918 1063 CB ALA 63 B 16.495 9.711 -1.407 1065 CA ALA 64 B 15.993 6.793 -0.077 1066 CA ALA 63 B 16.495 9.711 -1.407 1066 CA ALA 64 B 15.993 6.793 -0.077 1066 CA ALA 64 B 15.993 6.793 -0.077 1066 CA ALA 64 B 15.993 6.793 -0.077 1066 CA ALA 64 B 15.695 5.679 0.866 1069 N TYR 65 B 12.454 8.570 1.709 1071 C TYR 65 B 12.454 8.570 1.709 1077 C TYR 65 B 12.454 8.570 1.709 1077 C TYR 65 B 12.454 8.570 1.709 1077 C TYR 65 B 12.401 1.505 0.984 1076 CD2 TYR 65 B 12.454 8.570 1.709 1077 C TYR 65 B 12.401 1.505 0.984 1076 CD2 TYR 65 B 12.540 11.505 0.984 1076 CD2 TYR 65 B 12.540 11.505 0.984	A	tom	Res	duc	Chain	X	Y	Z
1037 O								
1038   CB   GLU   60   B   28.745   9.151   -0.867   1039   CG   GLU   60   B   27.995   10.260   -0.117   1040   CD   GLU   60   B   27.559   8.834   1.775   1042   OE2   GLU   60   B   26.329   10.634   1.518   1043   N   GLY   61   B   25.552   8.049   -1.395   1044   CA   GLY   61   B   24.342   7.623   -0.733   1045   C   GLY   61   B   23.253   8.252   -1.519   1046   O   GLY   61   B   23.535   8.769   -2.595   1047   N   TYR   62   B   22.021   8.246   -1.038   1048   CA   TYR   62   B   22.021   8.246   -1.038   1049   C   TYR   62   B   20.934   8.777   -1.830   1049   C   TYR   62   B   19.767   9.046   -0.932   1050   O   TYR   62   B   19.767   9.046   -0.932   1050   O   TYR   62   B   19.796   8.647   0.217   1051   CB   TYR   62   B   19.833   6.572   -2.370   1053   CD1   TYR   62   B   19.833   6.572   -2.370   1053   CD1   TYR   62   B   19.833   6.572   -2.370   1055   CE1   TYR   62   B   17.474   9.863   -0.693   1061   C   ALA   63   B   15.993   8.617   -1.981   1063   CB   ALA   64   B   15.293   6.791   -0.077   1066   CA   ALA   64   B   15.293   6.791   -0.066   1069   N   TYR   65   B   13.710   8.191   1.098   1071   C   TYR   65   B   13.710   8.191   1.098   1071   C   TYR   65   B   12.454   8.570   1.709   1071   C   TYR   65   B   12.540   11.505   0.984   1076   CD2   TYR   65   B   12.540	1036	С	GLU	60	В	26.735	7.734	-0.926
1039 CG GLU 60 B 27.995 10.280 -0.117 1040 CD GLU 60 B 27.231 9.868 1.144 1041 OE1 GLU 60 B 27.559 8.834 1.775 1042 OE2 GLU 60 B 26.329 10.634 1.518 1043 N GLY 61 B 25.552 8.049 -1.395 1044 CA GLY 61 B 23.555 8.769 -2.595 1046 O GLY 61 B 23.555 8.769 -2.595 1047 N TYR 62 B 22.021 8.246 -1.038 1048 CA TYR 62 B 20.934 8.777 -1.830 1049 C TYR 62 B 19.767 9.046 -0.932 1050 O TYR 62 B 19.767 9.046 -0.932 1050 C TYR 62 B 19.796 8.647 0.217 1051 CB TYR 62 B 19.833 6.572 -2.370 1053 CD1 TYR 62 B 20.425 5.395 -1.917 1054 CD2 TYR 62 B 19.624 4.316 -1.563 1056 CE2 TYR 62 B 19.624 4.316 -1.563 1056 CE2 TYR 62 B 17.592 5.644 -2.132 1059 N ALA 63 B 18.720 9.711 -1.407 1060 CA ALA 63 B 17.474 9.863 -0.693 1061 C ALA 63 B 17.474 9.863 -0.693 1062 O ALA 63 B 16.889 11.151 -1.155 1064 N ALA 64 B 16.262 7.847 0.047 1066 C ALA 63 B 15.903 8.617 -1.981 1067 O ALA 64 B 15.293 6.793 -0.077 1066 C ALA 65 B 12.454 8.570 1.709 1071 C TYR 65 B 12.454 8.570 1.709	1037	0	GLU					0.131
1040   CD   GLU   60   B   27.231   9.868   1.144     1041   OE1   GLU   60   B   27.555   8.834   1.775     1042   OE2   GLU   60   B   26.329   10.634   1.515     1043   N   GLY   61   B   25.552   8.049   -1.395     1044   CA   GLY   61   B   24.342   7.623   -0.733     1045   C   GLY   61   B   23.253   8.252   -1.519     1046   O   GLY   61   B   23.253   8.252   -1.519     1047   N   TYR   62   B   22.021   8.246   -1.038     1048   CA   TYR   62   B   22.021   8.246   -1.038     1049   C   TYR   62   B   20.934   8.777   -1.830     1049   C   TYR   62   B   19.767   9.046   -0.932     1050   O   TYR   62   B   19.767   9.046   -0.932     1050   O   TYR   62   B   19.796   8.647   0.217     1051   CB   TYR   62   B   20.608   7.745   -2.889     1052   CG   TYR   62   B   20.425   5.395   -1.917     1053   CD1   TYR   62   B   20.425   5.395   -1.917     1054   CD2   TYR   62   B   19.833   6.572   -2.370     1055   CE1   TYR   62   B   19.624   4.316   -1.563     1056   CE2   TYR   62   B   19.624   4.316   -1.563     1056   CE2   TYR   62   B   17.692   5.644   -2.132     1057   CZ   TYR   62   B   17.341   3.468   -1.352     1059   N   ALA   63   B   18.249   4.475   -1.681     1060   CA   ALA   63   B   18.720   9.711   -1.407     1060   CA   ALA   63   B   18.720   9.711   -1.407     1060   CA   ALA   63   B   15.903   8.617   -1.981     1063   CB   ALA   63   B   16.495   8.722   -0.918     1064   N   ALA   64   B   15.293   6.793   -0.077     1065   CA   ALA   64   B   15.293   6.793   -0.077     1066   C   ALA   64   B   15.293   6.793   -0.077     1067   O   ALA   64   B   15.695   5.679   0.666     1069   N   TYR   65   B   12.454   8.570   1.709     1071   C   TYR   65   B   12.454   8.570   1.709     1071   C   TYR   65   B   12.454   8.570   1.709     1073   CB   TYR   65   B   12.454   8.570   1.706     1074   CG   TYR   65   B   12.540   11.505   0.984     1075   CD1   TYR   65   B   12.540   11.505   0.984     1076   CD2   TYR   65   B   12.570   11.575   -1.721     1077   CE1   TYR   65	103B	CB	GLU				_	
1041 OE1 GLU 60 B 27.559 8.834 1.775 1042 OE2 GLU 60 B 26.329 10.634 1.518 1043 N GLY 61 B 25.552 8.049 -1.395 1044 CA GLY 61 B 23.555 8.252 -1.519 1046 C GLY 61 B 23.535 8.252 -1.519 1046 O GLY 61 B 23.535 8.769 -2.595 1047 N TYR 62 B 22.021 8.246 -1.038 1048 CA TYR 62 B 20.934 8.777 -1.830 1049 C TYR 62 B 19.767 9.046 -0.932 1050 O TYR 62 B 19.796 8.647 0.217 1051 CB TYR 62 B 20.608 7.745 -2.889 1052 CG TYR 62 B 19.833 6.572 -2.370 1053 CD1 TYR 62 B 20.425 5.395 -1.917 1054 CD2 TYR 62 B 19.624 4.316 -1.556 1056 CE2 TYR 62 B 19.624 4.316 -1.556 1056 CE2 TYR 62 B 19.624 4.316 -1.5681 1058 OB TYR 62 B 19.624 4.316 -1.5681 1059 N ALA 63 B 18.720 9.711 -1.407 1060 CA ALA 63 B 17.341 3.468 -1.352 1061 C ALA 63 B 16.485 8.722 -0.918 1062 O ALA 63 B 16.485 8.722 -0.918 1063 CB ALA 63 B 16.899 11.151 -1.155 1064 N ALA 64 B 15.293 6.793 -0.077 1065 CB ALA 64 B 15.293 6.793 -0.077 1066 C ALA 64 B 15.293 6.793 -0.077 1066 C ALA 64 B 15.695 5.679 0.866 1069 N TYR 65 B 12.454 8.570 1.709 1071 C TYR 65 B 12.454 8.570 1.709 1071 C TYR 65 B 12.454 8.570 1.709 1073 CB TYR 65 B 12.540 11.505 0.984 1076 CD2 TYR 65 B 12.570 11.505 0.984 1076 CD2 TYR 65 B 12.570 11.505 0.984 1077 CE1 TYR 65 B 12.571 11.575 -1.721	1039		GLU				10.280	-0.117
1042 OE2 GLU 60 B 26.329 10.634 1.51B 1043 N GLY 61 B 25.552 8.049 -1.395 1044 CA GLY 61 B 24.342 7.623 -0.733 1045 C GLY 61 B 23.253 8.252 -1.519 1046 O GLY 61 B 23.535 8.769 -2.595 1047 N TYR 62 B 22.021 8.246 -1.038 1048 CA TYR 62 B 20.934 8.777 -1.830 1049 C TYR 62 B 19.767 9.046 -0.932 1050 O TYR 62 B 19.766 8.647 0.217 1051 CB TYR 62 B 20.608 7.745 -2.889 1052 CG TYR 62 B 19.833 6.572 -2.370 1053 CD1 TYR 62 B 20.425 5.395 -1.917 1054 CD2 TYR 62 B 19.624 4.316 -1.563 1056 CE2 TYR 62 B 19.624 4.316 -1.563 1056 CE2 TYR 62 B 19.624 4.316 -1.563 1059 OH TYR 62 B 19.624 4.316 -1.563 1059 OH TYR 62 B 18.249 4.475 -1.681 1059 OH TYR 62 B 18.249 4.475 -1.681 1059 OH TYR 62 B 18.249 7.711 -1.407 1060 CA ALA 63 B 18.720 9.711 -1.407 1060 CA ALA 63 B 18.720 9.711 -1.407 1060 CA ALA 63 B 16.485 8.722 -0.918 1061 C ALA 63 B 16.485 8.722 -0.918 1062 O ALA 63 B 16.899 11.151 -1.155 1064 N ALA 64 B 15.293 6.793 -0.077 1065 CA ALA 64 B 15.293 6.793 -0.077 1066 C ALA 64 B 15.293 6.793 -0.077 1066 C ALA 64 B 15.293 6.793 -0.077 1067 O ALA 64 B 15.293 6.793 -0.077 1068 CB ALA 64 B 15.293 6.793 -0.077 1069 N TYR 65 B 13.710 8.191 1.098 1070 CA TYR 65 B 12.454 8.570 1.709 1071 C TYR 65 B 12.454 8.570 1.709 1071 C TYR 65 B 12.454 8.570 1.709 1074 CG TYR 65 B 12.508 10.513 0.168 1075 CD1 TYR 65 B 12.508 10.513 0.168 1075 CD2 TYR 65 B 12.571 11.751 -1.721								
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1047         N         TYR         62         B         22.021         8.246         -1.038           1048         CA         TYR         62         B         20.934         8.777         -1.830           1049         C         TYR         62         B         19.767         9.046         -0.932           1050         O         TYR         62         B         19.796         8.647         0.217           1051         CB         TYR         62         B         20.608         7.745         -2.889           1052         CG         TYR         62         B         19.833         6.572         -2.370           1053         CD1         TYR         62         B         19.824         4.316         -1.563           1054         CD2         TYR         62         B         19.624         4.316         -1.563           1055         CE1         TYR         62         B         17.692         5.644         -2.132           1057         CZ         TYR         62         B         17.341         3.468         -1.352           1059         N         ALA         63         B         17.								
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1054         CD2         TYR         62         B         18.481         6.692         -2.472           1055         CE1         TYR         62         B         19.624         4.316         -1.563           1056         CE2         TYR         62         B         17.692         5.644         -2.132           1057         CZ         TYR         62         B         18.249         4.475         -1.681           1058         OB         TYR         62         B         17.341         3.468         -1.352           1059         N         ALA         63         B         18.720         9.711         -1.407           1060         CA         ALA         63         B         17.474         9.863         -0.693           1061         C         ALA         63         B         16.485         8.722         -0.918           1062         O         ALA         63         B         15.903         8.617         -1.981           1063         CB         ALA         63         B         16.899         11.151         -1.155           1064         N         ALA         64         B         1								
1055         CE1         TYR         62         B         19.624         4.316         -1.563           1056         CE2         TYR         62         B         17.692         5.644         -2.132           1057         CZ         TYR         62         B         18.249         4.475         -1.681           1058         OH         TYR         62         B         17.341         3.468         -1.352           1059         N         ALA         63         B         18.720         9.711         -1.407           1060         CA         ALA         63         B         17.474         9.863         -0.693           1061         C         ALA         63         B         16.485         8.722         -0.918           1062         O         ALA         63         B         15.903         8.617         -1.981           1063         CB         ALA         63         B         16.899         11.151         -1.155           1064         N         ALA         64         B         15.293         6.793         -0.077           1065         CA         ALA         64         B         12								
1056         CE2         TYR         62         B         17.692         5.644         -2.132           1057         CZ         TYR         62         B         18.249         4.475         -1.681           1058         OH         TYR         62         B         17.341         3.468         -1.352           1059         N         ALA         63         B         18.720         9.711         -1.407           1060         CA         ALA         63         B         17.474         9.863         -0.693           1061         C         ALA         63         B         16.485         8.722         -0.918           1062         O         ALA         63         B         15.903         8.617         -1.981           1063         CB         ALA         63         B         16.899         11.151         -1.155           1064         N         ALA         64         B         15.293         6.793         -0.077           1065         CA         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         12.85								
1057         CZ         TYR         62         B         18.249         4.475         -1.681           1058         OH         TYR         62         B         17.341         3.468         -1.352           1059         N         ALA         63         B         18.720         9.711         -1.407           1060         CA         ALA         63         B         17.474         9.863         -0.693           1061         C         ALA         63         B         16.485         8.722         -0.918           1062         O         ALA         63         B         15.903         8.617         -1.981           1063         CB         ALA         63         B         16.899         11.151         -1.155           1064         N         ALA         64         B         16.262         7.847         0.047           1065         CA         ALA         64         B         15.293         6.793         -0.077           1066         C         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         15.695 </td <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>								
1058         OB         TYR         62         B         17.341         3.468         -1.352           1059         N         ALA         63         B         18.720         9.711         -1.407           1060         CA         ALA         63         B         17.474         9.863         -0.693           1061         C         ALA         63         B         16.485         8.722         -0.918           1062         O         ALA         63         B         15.903         8.617         -1.981           1063         CB         ALA         63         B         16.899         11.151         -1.155           1064         N         ALA         64         B         16.262         7.847         0.047           1065         CA         ALA         64         B         15.293         6.793         -0.077           1066         C         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         12.851         6.701         -0.265           1068         CB         ALA         64         B         15.695 </td <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>								
1059         N         ALA         63         B         18.720         9.711         -1.407           1060         CA         ALA         63         B         17.474         9.863         -0.693           1061         C         ALA         63         B         16.485         8.722         -0.918           1062         O         ALA         63         B         15.903         8.617         -1.981           1063         CB         ALA         63         B         16.899         11.151         -1.155           1064         N         ALA         64         B         16.262         7.847         0.047           1065         CA         ALA         64         B         15.293         6.793         -0.077           1066         C         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         12.851         6.701         -0.265           1068         CB         ALA         64         B         15.695         5.679         0.866           1069         N         TYR         65         B         12.454 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>								
1060         CA         ALA         63         B         17.474         9.863         -0.693           1061         C         ALA         63         B         16.485         8.722         -0.918           1062         O         ALA         63         B         15.903         8.617         -1.981           1063         CB         ALA         63         B         16.899         11.151         -1.155           1064         N         ALA         64         B         16.262         7.847         0.047           1065         CA         ALA         64         B         15.293         6.793         -0.077           1066         C         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         12.851         6.701         -0.265           1068         CB         ALA         64         B         15.695         5.679         0.866           1069         N         TYR         65         B         13.710         8.191         1.098           1070         CA         TYR         65         B         12.454 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>								
1061         C         ALA         63         B         16.485         8.722         -0.918           1062         O         ALA         63         B         15.903         8.617         -1.981           1063         CB         ALA         63         B         16.899         11.151         -1.155           1064         N         ALA         64         B         16.262         7.847         0.047           1065         CA         ALA         64         B         15.293         6.793         -0.077           1066         C         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         12.851         6.701         -0.265           1068         CB         ALA         64         B         15.695         5.679         0.866           1069         N         TYR         65         B         13.710         8.191         1.098           1070         CA         TYR         65         B         12.454         8.570         1.709           1071         C         TYR         65         B         12.101								
1062         O         ALA         63         B         15.903         8.617         -1.981           1063         CB         ALA         63         B         16.899         11.151         -1.155           1064         N         ALA         64         B         16.262         7.847         0.047           1065         CA         ALA         64         B         15.293         6.793         -0.077           1066         C         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         12.851         6.701         -0.265           1068         CB         ALA         64         B         15.695         5.679         0.866           1069         N         TYR         65         B         13.710         8.191         1.098           1070         CA         TYR         65         B         12.454         8.570         1.709           1071         C         TYR         65         B         11.742         7.437         2.314           1072         O         TYR         65         B         12.101								
1063         CB         ALA         63         B         16.899         11.151         -1.155           1064         N         ALA         64         B         16.262         7.847         0.047           1065         CA         ALA         64         B         15.293         6.793         -0.077           1066         C         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         12.851         6.701         -0.265           1068         CB         ALA         64         B         15.695         5.679         0.866           1069         N         TYR         65         B         13.710         8.191         1.098           1070         CA         TYR         65         B         12.454         8.570         1.709           1071         C         TYR         65         B         11.742         7.437         2.314           1072         O         TYR         65         B         12.101         6.294         2.178           1073         CB         TYR         65         B         11.531								
1064         N         ALA         64         B         16.262         7.847         0.047           1065         CA         ALA         64         B         15.293         6.793         -0.077           1066         C         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         12.851         6.701         -0.265           1068         CB         ALA         64         B         15.695         5.679         0.866           1069         N         TYR         65         B         13.710         8.191         1.098           1070         CA         TYR         65         B         12.454         8.570         1.709           1071         C         TYR         65         B         11.742         7.437         2.314           1072         O         TYR         65         B         12.101         6.294         2.178           1073         CB         TYR         65         B         11.531         9.231         0.716           1074         CG         TYR         65         B         12.088								
1065         CA         ALA         64         B         15.293         6.793         -0.077           1066         C         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         12.851         6.701         -0.265           1068         CB         ALA         64         B         15.695         5.679         0.866           1069         N         TYR         65         B         13.710         8.191         1.098           1070         CA         TYR         65         B         12.454         8.570         1.709           1071         C         TYR         65         B         11.742         7.437         2.314           1072         O         TYR         65         B         12.101         6.294         2.178           1073         CB         TYR         65         B         11.531         9.231         0.716           1074         CG         TYR         65         B         12.088         10.513         0.168           1075         CD1         TYR         65         B         12.170								
1066         C         ALA         64         B         13.847         7.203         0.227           1067         O         ALA         64         B         12.851         6.701         -0.265           1068         CB         ALA         64         B         15.695         5.679         0.866           1069         N         TYR         65         B         13.710         8.191         1.098           1070         CA         TYR         65         B         12.454         8.570         1.709           1071         C         TYR         65         B         11.742         7.437         2.314           1072         O         TYR         65         B         12.101         6.294         2.178           1073         CB         TYR         65         B         11.531         9.231         0.716           1074         CG         TYR         65         B         12.088         10.513         0.168           1075         CD1         TYR         65         B         12.540         11.505         0.984           1076         CD2         TYR         65         B         12.170					•			
1067         O         ALA         64         B         12.851         6.701         -0.265           1068         CB         ALA         64         B         15.695         5.679         0.866           1069         N         TYR         65         B         13.710         8.191         1.098           1070         CA         TYR         65         B         12.454         8.570         1.709           1071         C         TYR         65         B         11.742         7.437         2.314           1072         O         TYR         65         B         12.101         6.294         2.178           1073         CB         TYR         65         B         11.531         9.231         0.716           1074         CG         TYR         65         B         12.088         10.513         0.168           1075         CD1         TYR         65         B         12.540         11.505         0.984           1076         CD2         TYR         65         B         12.170         10.655         -1.176           1077         CE1         TYR         65         B         13.147<								
1068         CB         ALA         64         B         15.695         5.679         0.866           1069         N         TYR         65         B         13.710         8.191         1.098           1070         CA         TYR         65         B         12.454         8.570         1.709           1071         C         TYR         65         B         11.742         7.437         2.314           1072         O         TYR         65         B         12.101         6.294         2.178           1073         CB         TYR         65         B         11.531         9.231         0.716           1074         CG         TYR         65         B         12.088         10.513         0.168           1075         CD1         TYR         65         B         12.540         11.505         0.984           1076         CD2         TYR         65         B         12.170         10.655         -1.176           1077         CE1         TYR         65         B         13.147         12.614         0.454           1078         CE2         TYR         65         B         12.77								
1069         N         TYR         65         B         13.710         8.191         1.098           1070         CA         TYR         65         B         12.454         8.570         1.709           1071         C         TYR         65         B         11.742         7.437         2.314           1072         O         TYR         65         B         12.101         6.294         2.178           1073         CB         TYR         65         B         11.531         9.231         0.716           1074         CG         TYR         65         B         12.088         10.513         0.168           1075         CD1         TYR         65         B         12.540         11.505         0.984           1076         CD2         TYR         65         B         12.170         10.655         -1.176           1077         CE1         TYR         65         B         13.147         12.614         0.454           1078         CE2         TYR         65         B         12.771         11.751         -1.721								
1070         CA         TYR         65         B         12.454         8.570         1.709           1071         C         TYR         65         B         11.742         7.437         2.314           1072         O         TYR         65         B         12.101         6.294         2.178           1073         CB         TYR         65         B         11.531         9.231         0.716           1074         CG         TYR         65         B         12.088         10.513         0.168           1075         CD1         TYR         65         B         12.540         11.505         0.984           1076         CD2         TYR         65         B         12.170         10.655         -1.176           1077         CE1         TYR         65         B         13.147         12.614         0.454           1078         CE2         TYR         65         B         12.771         11.751         -1.721								
1071         C         TYR         65         B         11.742         7.437         2.314           1072         O         TYR         65         B         12.101         6.294         2.178           1073         CB         TYR         65         B         11.531         9.231         0.716           1074         CG         TYR         65         B         12.088         10.513         0.168           1075         CD1         TYR         65         B         12.540         11.505         0.984           1076         CD2         TYR         65         B         12.170         10.655         -1.176           1077         CE1         TYR         65         B         13.147         12.614         0.454           1078         CE2         TYR         65         B         12.771         11.751         -1.721								
1072     O     TYR     65     B     12.101     6.294     2.178       1073     CB     TYR     65     B     11.531     9.231     0.716       1074     CG     TYR     65     B     12.088     10.513     0.168       1075     CD1     TYR     65     B     12.540     11.505     0.984       1076     CD2     TYR     65     B     12.170     10.655     -1.176       1077     CE1     TYR     65     B     13.147     12.614     0.454       1078     CE2     TYR     65     B     12.771     11.751     -1.721								
1073         CB         TYR         65         B         11.531         9.231         0.716           1074         CG         TYR         65         B         12.088         10.513         0.168           1075         CD1         TYR         65         B         12.540         11.505         0.984           1076         CD2         TYR         65         B         12.170         10.655         -1.176           1077         CE1         TYR         65         B         13.147         12.614         0.454           1078         CE2         TYR         65         B         12.771         11.751         -1.721								
1074     CG     TYR     65     B     12.088     10.513     0.168       1075     CD1     TYR     65     B     12.540     11.505     0.984       1076     CD2     TYR     65     B     12.170     10.655     -1.176       1077     CE1     TYR     65     B     13.147     12.614     0.454       1078     CE2     TYR     65     B     12.771     11.751     -1.721								
1075     CD1     TYR     65     B     12.540     11.505     0.984       1076     CD2     TYR     65     B     12.170     10.655     -1.176       1077     CE1     TYR     65     B     13.147     12.614     0.454       1078     CE2     TYR     65     B     12.771     11.751     -1.721								
1076     CD2     TYR     65     B     12.170     10.655     -1.176       1077     CE1     TYR     65     B     13.147     12.614     0.454       1078     CE2     TYR     65     B     12.771     11.751     -1.721								
1077 CE1     TYR 65     B     13.147     12.614     0.454       1078 CE2     TYR 65     B     12.771     11.751     -1.721								
1078 CE2 TYR 65 B 12.771 11.751 -1.721								
1070 C7 TVD 65 to 12 707 17 701 0 000	1078	CE2		65	B	13.282	12.701	-1.721
1079 CZ TYR 65 B 13.282 12.701 -0.902 1080 OH TYR 65 B 13.974 13.762 -1.462								

Fig. 15.25

A	tom	Resi	duc	Chain	x	<u>Y</u>	z
1081	N	TYR	66	В	10.697	7.722	3.031
1082	CA	TYR	66	В	9.852	6.661	3.471
1083	С	TYR	66	В	8.541	7.291	3.740
1084	0	TYR	66	В	8.435	8.508	3.867
1085	СВ	TYR	66	В	10.389	5.996	4.739
1086	CG	TYR	66	В	10.325	6.790	6.014
1087	CD1	TYR	66	В	11.182	7.824	6.193
1088	CD2	TYR	66	В	9.448	6.427	7.006
1089	CE1	TYR	66	В	11.162	8.514	7.369
1090	CE2	TYR	66	В	9.415	7.105	8.185
1091	CZ	TYR	66	В	10.273	8.151	8.343
1092	OH	TYR	66	В	10.254	8.912	9.507
1093	N	CYS	67	В	7.523	6.461	3.838
1094	CA	CYS	67	В	6.157	6.944	3.974
1095	С	CYS	67	В	5.600	6.711	5.382
1096	0	CYS	67	В	5.660	5.589	5.909
1097	CB	CYS	67	В	5.210	6.219	3.078
1098	SG	CYS	67	В .	5.398	6.240	1.331
1099	N	GLU	68	В	5.026	7.712	6.056
1100	CA	GLU	68	В	4.554	7.337	7.342
1101	C	GLU	68	В	3.507	8.263	7.751
1102	0	GLU	68	В	3.670	9.457	7.509
1103	CB	GLU	68	В	5.639	7.411	8.360
1104	CG	GLU	68	В	5. <b>32</b> 2	6.494	9.510
1105	CD	GLU	68	В	6.024	7.006	10.718
1106	OE1	GLU	68	В	6.127	6.266	11.697
1107	OE2	GLU	68	В	6.475	8.154	10.689
1108	N	GLY	69	В	2.450	7.734	8.369
1109	CA	GLY	69	В	1.458	8.577	9.002
1110	C	GLY	69	В	0.123	7.873	8.939
1111	0	GLY	69	В	0.019	6.722	8.517
1112	N	GLU	70	В	-0.962	8.499	9.336
1113	CA	GLU	70	В	-2.120	7.685	9.616
1114	С	GLU	70	В	-3.133	7.672	8.511
1115	0	GLU	70	В	-3.372	8.697	7.898
1116	CB	GLU	70	В	-2.769	8.180	10.882
1117	CG	GLU	70	В	-2.975	6.949	11.748
1118	CD	GLU	70	В	-4.167	7.206	12.630
1119	OE1	GLU	70	В	-4.866	6.234	12.949
1120	OE2	GLU	70	В	-4.412	8.381	12.984
1121	N	CYS	71	В	-3.757	6.544	8.233
1122	CA	CYS	71	В	-4.817	6.538	7.262
1123	C.	CYS	71	В	-6.205	6.516	7.840
1124	0	CYS	71	В	-6.730	5.505	8.245
1125	CB	CYS	71	В	-4.553	5.350	6.377

Fig. 15.26

A	tom	Resi	duc	Chain	x	<u>Y</u>	Z
1126	SG	CYS	71	. В	-3.363	5.857	5.112
1127	N	ALA	72	В	-6.909	7.619	7.924
1128	CA	ALA	72	В	-8.091	7.666	8.737
1129	С	ALA	72	В	-0.935	8.777	8.184
1130	0	ALA	72	В	-8.438	9.590	7.418
1131	CB	ALA	72	В	-7.646	7.939	10.149
1132	N	PHE	73	В	-10.225	8.934	8.471
1133	CA	PHE	73	В	-10.963	9.998	7.782
1134	C	PHE	73	B	-10.507	11.313	8.342
1135	0	PHE	73	В	-10.291	11.328	9.544
1136	CB	PHE	73	В	-12.492	9.874	7.997
1137	CG	PHE	73	В	-13.023	8.690	7.235
1138	CD1	PHE	73	В	-13.009	8.688	5.873
1139	CD2	PHE	73	В	-13.514	7.616	7.919
1140	CE1	PHE	73	В	-13.473	7.602	5.190
1141	CE2	PHE	73	В	-13.992	6.528	7.214
1142	CZ	PHE	73	В	-13.969	6.518	5.854
1143	N	PRO	74	В	-10.337	12.410	7.712
1144	CA	PRO	74	В	-10.748	12.685	6.357
1145	С	PRO	74	В	-9.754	12.033	5.502
1146	0	PRO	74	В	-8.656	12.529	5.443
1147	CB	PRO	74	В	-10.712	14.168	6.197
1148	CG	PRO	74	В	-10.432	14.652	7.611
1149	CD	PRO	74	В	-9.662	13.546	8.313
1150	N	LEU	75	В	-10.018	10.938	4.799
1151	CA	LEU	75	В	-9.091	10.355	3.845
1152	С	LEU	75	В	-9.061	11.400	2.779
1153	0	LEU	75	В	-9.752	11.427	1.772
1154	CB	TEA	75	В	-9.674	9.034	3.355
1155	CG	TEA	75	В	-8.684	7.913	3.407
1156	CD1	LEU	75	В	-7.689	8.120	4.525
1157	CD2	LEU	75	В	-9.442	6.631	3.626
1159	N	asn	76	В	-8.181	12.333	3.059
1159	CA	asn	76	В	-8.098	13.558	2.311
1160	C	ASN	76	В	-7.517	13.432	0.887
1161	0	asn	76	В	-6.525	14.065	0.596
1162	CB	ASN	76	В	-7.310	14.402	3.255
1163	CG	asn	76	В	-7.466	15.837	2.965
1164	OD1	ASN	76	В	-7.449	16.655	3.894
1165	ND2	ASN	76	В	-7.521	16.219	1.674
1166	N	SER	77	В	-8.021	12.652	-0.088
1167	CA	SER	77	В	-7.673	12.747	-1.523
1168	C	SER	77	В	-6.200	12.824	-1.891
1169	0	SER	77	В	-5.587	11.803	-2.261
1170	CB	SER	77	В	-8.354	13.964	-2.171

Fig. 15.27

A	tom	Resi	idue	Chain	<u>x</u>	Y	Z
1171	OG	SER	77	В	-7.957	15.177	-1.532
1172	И	TYR	78	В	-5.531	13.986	-1.819
1173	CA	TYR	78	<b>B</b> .	-4.098	13.882	-1.965
1174	С	TYR	78	В	-3.490	13.159	-0.764
1175	0	TYR	78	В	-2.585	13.733	-0.185
1176	CB	TYR	78	В	-3.438	15.241	-2.045
1177	CG	TYR	78	В	-2.916	15.632	-3.429
1178	CD1	TYR	78	В	-1.897	16.590	-3.392
1179	CD2	TYR	78	В	-3.617	15.385	-4.630
1180	CE1	TYR	78	В	-1.737	17.490	-4.438
1181	CE2	TYR	78	В	-3.456	16.292	-5.692
1182	CZ	TYR	78	В	-2.610	17.403	-5.516
1183	OH	TYR	78	В	-2.715	18.527	-6.319
1184	N	MET	79	В	-3.978	11.959	-0.393
1185	CA	MET	79	В	-3.388	10.933	0.448
1186	С	MET	79	В	-3.689	9.603	-0.251
1187	0	MET	79	B	-3.734	8.505	0.285
1188	CB	MET	79	В	-4.031	10.775	1.776
1189	CG	MET	79	В	-4.050	11.928	2.741
1190	SD	MET	79	В	-4.737	11.135	4.215
1191	CE	MET	79	В	-4.910	12.550	5.255
1192	N	ASN	80	В	-3.926	9.699	-1.544
1193	CA	ASN	80	В	-4.199	8.581	-2.401
1194	С	ASN	80	В	-5.079	7.439	-1.962
1195	0	ASN	80	В	-4.924	6.364	-2.490
1196	CB	ASN	80	В .	-2.883	7.967	-2.894
1197	CG	ASN	80	В	-3.155	7.090	-4.121
1198	OD1	ASN	80	В	-3.135	5.844	-4.162
1199	ND2	ASN	80	В	-3.543	7.769	-5.1 <sub>.</sub> 98
1200	N	ALA	81	В	-6.051	7.407	-1.058
1201	CA	ALA	81	В	-7.299	6.757	-1.424
1202	С	ALA	81	В	-7.486	5.344	-1.928
1203	0	ALA	81	В	-8.004	4.449	~1.302
1204	CB	ALA	81	В	-8.004	7.624	-2.452
1205	N	THR	82	В	-7.106	4.970	-3.106
1206	CA	THR	82	В	-7.875	3.957	-3.819
1207	C	THR	82	В	-9.402	3.970	-3.655
1208	0	THR	82	В	-10.075	4.641	-4.444
1209	CB	THR	82	B	-7.505	2.477	-3.539
1210	OG1	THR	82	В	-7.534	2.217	-2.163
1211	CG2	THR	82	В	-6.211	2.155	-4.224
1212	N	ASN	83	В	-10.043	3.286	-2.697
1213	CA	ASN	83	В	-11.462	2.977	-2.877
1214	С	ASN	83	В	-11.670	1.861	-1.963
1215	0	ASN	83	В	-12.472	1.858	-1.070

Fig. 15.28

A	tom	Resi	duc	Chain	X	<u>Y</u>	<u>Z</u>
1216	CB	ASN	83	В	-11.835	2.404	-4.195
1217	CG	ASN	83	В	-12.710	3.302	-4.983
1218	OD1	ASN	83	В	-13.343	2.872	-5.931
1219	ND2	ASN	83	В	-12.741	4.594	-4.715
1220	N	HIS	84	В	-10.950	0.775	-2.095
1221	CA	HIS	84	В	-10.823	-0.192	-1.024
1222	C	HIS	84	В	-10.284	0.531	0.167
1223	o	HIS	84	В	-10.623	0.097	1.245
1224	СВ	HIS	84	В	-9.851	-1.284	-1.397
1225	CG	HIS	84	В	-9.778	-2.364	-0.370
1226	ND1	HIS	84	В	-9.014	-2.402	0.711
1227	CD2	HIS	84	В	-10.397	-3.578	-0.489
1228	CE1	HIS	84	В	-9.153	-3.590	1.235
1229	NE2	HIS	84	В	-9.960	-4.294	0.507
1230	N	ALA	85	В	-9.464	1.589	0.169
1231	CA	ALA	85	В	-9.158	2.136	1.455
1232	C	ALA	85	В	-10.314	2.915	1.965
1233	0	ALA	85	В	-10.388	3.113	3.155
1234	СВ	ALA	85	В	-7.958	3.026	1.398
1235	N	ILE	86	В	-11.298	3.418	1.242
1236	CA	ILE	86	В	-12.366	4.096	1.919
1237	С	ILE	86	В	-13.219	2.964	2.461
1238	0	ILE	86	В	-13.597	3.036	3.590
1239	СВ	ILE	86	В	-13.058	4.975	0.897
1240	CG1	ILE	В6	В	-12.035	5.957	0.299
1241	CG2	ILE	86	В	-14.230	5.699	1.565
1242	CD1	ILE	В6	В	-12.568	7.113	-0.547
1243	N	VAL	87	В	-13.596	1.857	1.829
1244	CA	VAL	87	В	-14.315	0.785	2.493
1245	С	VAL	87	В	-13.560	0.425	3.744
1246	ō	VAL	87	В	-14.049	0.656	4.828
1247	СВ	VAL	87	В	-14.432	-0.429	1.553
1248	CG1	VAL	87	В	-14.652	-1.712	2.308
1249	CG2	VAL	87	В	-15.617	-0.236	0.637
1250	N	GLN	88	В	-12.358	-0.137	3.734
1251	CA	GLN	88	В	-11.706	-0.524	4.975
1252	С	GLN	88	В	-11.666	0.574	6.022
1253	0	GLN	88	В	-11.943	0.280	7.170
1254	СВ	GLN	88	В	-10.295	-0.941	4.697
1255	CG	GLN	88	В	-9.864	-2.322	5.125
1256	CD	GLN	88	В	-8.398	-2.237	5.020
1257	OE1	GLN	88	В	-7.618	-2.317	5.944
1258	NE2	GLN	88	В	-7.922	-1.853	3.852
1259	N	THR	89	В	-11.347	1.833	5.754
	CA	THR	89	В	-11.401	2.802	6.790

Fig. 15.29

A1	tom	Resi	duc	Chain	<u>x</u>	Υ	<u>z</u>
1261	С	THR	89	В	-12.805	2.953	7.298
1262	0	THR	89	В	-12.966	3.090	8.499
1263	CB	THR	89	B	-10.923	4.081	6.270
1264	OG1	THR	89	В	-9.628	3.792	5.879
1265	CG2	THR	89	В	-10.741	5.183	7.270
1266	N	LEU	90	В	-13.888	2.943	6.506
1267	CA	LEU	90	В	-15.260	2.885	7.003
1268	С	LEU	90	В	-15.550	1.649	7.814
1269	0	LEU	90	В	-16.030	1.727	8.929
1270	CB	LEU	90	В	-16.220	2.893	5.853
1271	CG	LEU	90	В	-17.670	3.011	6.158
1272	CD1	LEU	90	В	-17.933	4.270	6.954
1273	CD2	LEU	90	В	-18.406	3.003	4.837
1274	N	VAL	91	В	-15.296	0.449	7.333
1275	CA	VAL	91	В	-15.459	-0.745	8.132
1276	С	VAL	91	В	-14.637	-0.671	9.387
1277	0	VAL	91	В	-14.959	-1.278	10.398
1278	CB	VAL	91	В	-15.049	-1.923	7.308
1279	CG1	VAL	91	В	-14.917	-3.200	8.111
1280	CG2	VAL	91	В	-16.122	-2.071	6.250
1281	N	HIS	92	В	-13.533	0.068	9.434
1282	ÇA	HIS	92	В	-12.818	0.180	10.702
1283	С	BIS	92	В	-13.719	0.784	11.705
1284	0	HIS	92	В	-13.859	0.310	12.797
1285	CB	HIS	92	В	-11.577	1.095	10.619
1286	CG	HIS	92	В	-10.675	0.908	11.812
1287	ND1	HIS	92	В	-10.106	1.714	12.701
1288	CD2	HIS	92	В	-10.314	-0.348	12.142
1289	CEl	HIS	92	В	-9.454	0.950	13.527
1290	NE2	HIS	92	В	-9.602	-0.300	13.197
1291	N	PHE	93	В	-14.364	1.854	11.358
1292	CA	PHE	93	В	-15.148	2.645	12.248
1293	C	PHE	93	В	-16.394	1.899	12.634
1294	ō	PHE	93	В	-16.782	1.756	13.777
1295	CB	PHE	93	В	-15.365	3.894	11.461
1296	CG	PHE	93	В	-16.515	4.726	11.911
1297	CD1	PHE	93	В	-17.812	4.330	11.649
1298	CD2	PHE	93	В	-16.248	5.977	12.448
1299	CE1	PHE	93	В	-18.832	5.210	11.890
1300	CE2	PHE	93	В	-17.282	6.860	12.670
1301	CZ	PHE	93	В	-18.564	6.467	12.386
1302	N	ILE	94	В	-17.172	1.345	11.730
1303	CA	ILE	94	В	-18.251	0.460	12.158
1304	c	ILE	94	В	-17.714	-0.589	13.105
1305	o	ILE	94	В	-18.482	-1.254	13.758

Fig. 15.30

At	om_	Res	idue	Chain	x	<u>Y</u>	z
				_			
1306	CB	ILE	94	В	-18.808	-0.124	10.880
1307	CG1	ILE	94	В	-19.580	1.028	10.342
1308	CG2	ILE	94	В	-19.648	-1.375	10.977
1309	CD1	ILE	94	В	-19.872	0.740	8.877
1310	N	asn	95	B	-16.410	-0.839	13.277
1311	CA	ASN	95	В	-16.013	-2.092	13.873
1312	C.	ASN	95	В	-14.529	-2.322	14.157
1313	0	asn	95	В	-13.858	-3.248	13.673
1314	CB	asn	95	В	-16.495	-3.208	12.993
1315	CG	ASN	95	В	-16.663	-4.358	13.942
1316	OD1	ASN	95	В	-16.021	-4.431	15.007
1317	ND2	ASN	95	В	-17.547	-5.309	13.564
1318	N	PRO	96	В	-13.981	-1.497	14.982
1319	CA	PRO	96	В	-12.565	-1.248	15.087
1320	С	PRO	96	В	-11.697	-2.421	15.300
1321	0	PRO	96	В	-10.491	-2.366	15.067
1322	СВ	PRO	96	В	-12.426	-0.227	16.199
1323	CG	PRO	96	B	-13.819	0.347	16.282
1324	CD	PRO	96	В	-14.625	-0.944	16.151
1325	N	GLU	97	В	-12.338	-3.478	15.747
1326	CA	GLU	97	В	-11.625	-4.700	15.963
1327	С	GLU	97	В	-11.796	-5.472	14.691
1328	0	GLU	97	В	-10.808	-5.919	14.166
1329	CB	GLU	97	В	-12.209	<b>-5.474</b>	17.153
1330	CG	GLU	97	В	-11.332	-5.321	. 18.423
1331	CD	GLU	97	В	-11.552	-3.957	19.136
1332	OE1	GLU	97	В	-12.507	-3.902	19.971
1333	OE2	GLU	97	В	-10.781	-2.974	18.873
1334	N	THR	98	В	-12.921	-5.722	14.047
1335	CA	THR	98	В	-12.848	-6.579	12.862
1336	С	THR	98	В	-11.921	-6.174	11.682
1337	0	THR	98	В	-11.787	-7.035	10.815
1338	CB	THR	98	В	-14.289	-6.757	12.349
1339	OG1	THR	98	В	-14.841	-5.463	12.337
1340	CG2	THR	98	В	-15.226	-7.552	13.281
1341	N	VAL	99	В	-11.255	-5.008	11.488
1342	CA	VAL	99	В	-10.426	-4.763	10.307
1343	С	VAL	99	В	-9.447	-3.636	10.637
1344	0	VAL	99	В	-9.956	-2.741	11.289
1345	CB	VAL	99	В	-11.316	-4.375	9.112
1346	CG1	VAL	99	В	-11.405	-2.876	8.750
1347	CG2	VAL	99	В	-10.670	-5.076	7.950
1348	N	PRO	100	В	-8.179	-3.500	10.300
1349	CA	PRO	100	В	-7.213	-2.555	10.669
1350	С	PRO		В	-7.409	-1.258	10.007

Fig. 15.31

A	tom	Residue	Chain	X	Y	Z
1351	0	PRO 100	В	-8.381	-1.158	9.326
1352	CB	PRO 100	В	-5.890	-3.130	10.342
1353	CG	PRO 100	В	-6.120	-4.613	10.443
1354	CD	PRO 100	В	-7.484	-4.771	9.830
1355	N	LYS 101	В	-6.660	-0.169	10.060
1356	CA	LYS 101	В	-7.015	0.958	9.218
1357	С	LYS 101	В	-6.112	0.771	8.015
1358	0	LYS 101	В	-5.130	0.061	8.142
1359	CB	LYS 101	В	-6.671	2.300	9.855
1360	CG	LYS 101	В	-6.510	2.347	11.360
1361	CD	LYS 101	В	-5.813	3.616	11.889
1362	CE	LYS 101	В	-4.227	3.513	12.046
1363	NZ	LYS 101	B	-3.387	3.806	10.849
1364	N	PRO 102	В	-6.219	1.262	6.840
1365	CA	PRO 102	В	-5.321	0.916	5.769
1366	С	PRO 102	В	-3.970	1.433	6.090
1367	0	PRO 102	B	-3.853	2.346	6.892
1368	CB	PRO 102	В	-5.934	1.534	4.529
1369	ÇG	PRO 102	В	-6.576	2.741	5.074
1370	CD	PRO 102	В	-7.100	2.341	6.447
1371	N	CYS 103	В	-2.852	0.963	5.547
1372	CA	CYS 103	В	-1.653	1.665	5.922
1373	С	CYS 103	В	-0.993	2.294	4.757
1374	0	CYS 103	В	-1.354	2.179	3.600
1375	CB	CYS 103	В	-0.697	0.722	6.598
1376	SG	CYS 103	В	0.104	-0.064	5.231
1377	N	CYS 104	В	0.044	2.988	5.123
1378	CA	CYS 104	В	0.501	4.087	4.348
1379	С	CYS 104	В	1.771	3.676	3.652
1380	0	CYS 104	В	2.791	3.408	4.251
1381	CB	CYS 104	В	0.602	5.107	5.385
1382	SG	CYS 104	В	1.414	6.474	4.692
1383	N	ALA 105	В	1.749	3.613	2.347
1384	CA	ALA 105	В	2.637	2.781	1.614
1385	C	ALA 105	В	3.081	3.640	0.494
1386	0	ALA 105	В	2.412	4.644	0.296
1387	CB	ALA 105	B	1.841	1.604	1.154
1388	N	PRO 106	B	4.100	3.411	-0.258
1389	CA	PRO 106	В	4.400	4.197	-1.420
1390	С	PRO 106	В	3.597	3.811	-2.662
1391	0	PRO 106	B	3.311	2.672	-2.978
1392	CB	PRO 106	B	5.891	4.039	-1.557
1393	CG	PRO 106	В .	6.123	2.662	-1.081
1394	CD	PRO 106	В	5.349	2.826	0.199
1395	N	THR 107	В	3.246	4.864	-3.375

Fig. 15.32

A	tom	Residuc	Chain	<u> </u>	Y	Z
1206	G.	mun 107	-	2 221	4 005	
1396	CA C	THR 107 THR 107	B B	2.321	4.885	-4.464
1397	0	THR 107	В	3.250	4.827	-5.644
1398 1399	CB	THR 107	В	3.206 1.582	3.871 6.175	-6.411
1400	OG1	THR 107	B	0.518	5.644	-4.155 -3.377
1401	CG2	THR 107	В	1.134	7.066	-5.292
1402	N	GLN 108	₽.	4.151	5.763	-5.922
1403	CA	GLN 108	В	5.164	5.360	-6.846
1404	C	GLN 108	В	6.603	5.599	-6.478
1405	0	GLN 108	B	6.989	6.589	-5.880
1406	СВ	GLN 108	В′	4.712	6.001	-8.168
1407	CG	GLN 108	В	5.202	7.362	-8.639
1408	CD	GLN 108	В	4.487	8.380	-7.859
1409	OE1	GLN 108	В	3.911	8.140	-6.806
1410	NE2	GLN 108	В	4.552	9.578	-8.441
1411	N	LEU 109	В	7.351	4.579	-6.896
1412	CA	LEU 109	В	8.773	4.347	-6.622
1413	c	LEU 109	В	9.747	4.700	-7.774
1414	0	LEU 109	В	9.510	4.496	-8.963
1415	CB	LEU 109	В	8.945	2.848	-6.208
1416	CG	LEU 109	В	8.351	2.584	-4.813
1417	CD1	LEU 109	В	8.260	1.128	-4.489
1418	CD2	LEU 109	В	9.269	3.213	-3.792
1419	N	ASN 110	В	10.911	5.262	-7.494
1420	CA	ASN 110	В	11.823	5.551	-8.555
1421	С	ASN 110	В	13.077	4.792	-8.391
1422	0	ASN 110	В	13.238	4.036	-7.445
1423	CB	ASN 110	В	12.199	6.976	-8.586
1424	CG	ASN 110	В	11.025	7.786	-8.983
1425	OD1	ASN 110	В	11.064	9.002	-8.744
1426	ND2	ASN 110	В	9.942	7.206	-9.529
1427	N	ALA 111	В	14.021	4.976	-9.308
1428	CA	ALA 111	В	15.130	4.055	-9.447
1429	С	ALA 111	В	16.280	4.897	-9.088
1430	0	ALA 111	В	16.134	6.101	-9.021
1431	CB	ALA 111	В	15.249	3.543	-10.900
1432	N	ILE 112	В	17.436	4.324	-8.843
1433	CA	ILE 112	В	18.624	5.094	-8.502
1434	С	ILE 112	В	19.626	4.386	-9.369
1435	0	ILE 112	В	19.520	3.175	-9.473
1436	СВ	ILE 112	В	18.834	4.979	-6.944
1437	CG1	ILE 112	В	18.727	6.452	-6.591
1438	CG2	ILE 112	В	20.119	4.396	-6.342
1439	CD1	ILE 112	В	20.014	7.252	-6.922
1440	N	SER 113	В	20.575	5.052	-9.998

Fig. 15.33

A	om	Residue	Chain	x	Y	
			÷		4 222	44 445
1441	CA .	SER 113	B -	21.646		-10.635
1442	С	SER 113	B	22.926	4.454	-9.879
1443	0	SER 113	В	23.220	5.579	-9.520
1444	CB	SER 113	B	21.834		-12.034
1445	OG	SER 113	B	21.085		-12.909
1446	N	VAL 114	B	23.726	3.408	-9.592
1447	CA	VAL 114	В	25.005	3.645	-8.929
1448	C	VAL 114	В	26.085	3.074	-9.841
1449	0	VAL 114	В	25.914		-10.484
1450	CB	VAL 114	В	25.150	2.977	-7.496
1451	CG1	VAL 114	В	23.888	3.143	-6.703
1452	CG2	VAL 114	B	25.446	1.508	-7.575
1453	N	LEU 115	В	27.215	3.773	-9.911
1454	CA	LEU 115	В	28.417		-10.663
1455	С	LEU 115	В	29.406	2.624	-9.859
1456	0	LEU 115	В	29.857	3.095	-8.814
1457	CB	LEU 115	В	29.096		-11.145
1458	CG	LEU 115	В	30.445		-11.850
1459	CD1	LEU 115	В	30.271		-13.289
1460	CD2	LEU 115	В	31.054		-11.750
1461	N	TYR 116	В	29.854		-10.212
1462	CA	TYR 116	В	30.750	0.783	-9.287
1463	С	TYR 116	В	31.728	-0.199	-9.868
1464	0	TYR 116	В	31.498		-10.976
1465	CB	TYR 116	В	29.872	0.123	-8.278
1466	CG	TYR 116	В	29.144	-1.012	-B.907
1467	CD1	TYR 116	В	27.919	-0.788	
1468	CD2	TYR 116	В	29.728	-2.243	-B.856
1469	CEl	TYR 116	В	27.289		-10.101
1470	CE2	TYR 116	В	29.105	-3.280	-9.474
1471	CZ	TYR 116	В	27.925		-10.113
1472	OH	TYR 116	В	27.396		-10.879
1473	N	PHE 117	В	32.820	-0.583	-9.201
1474	CA	PHE 117	В	33.733	-1.520	-9.828
1475	С	PHE 117	В	33.168	-2.861	-9.521
1476	0	PHE 117	В	33.029	-3.159	-8.347
1477	CB	PHE 117	В	35.137	-1.590	-9.242
1478	CG	PHE 117	В	35.701	-0.236	-8.937
1479	CD1	PHE 117	В	35.512	0.315	-7.691
1480	CD2	PHE 117	В	36.392	0.459	-9.903
1481	CE1	PHE 117	B	36.002	1.579	
1482	CE2	PHE 117	В	36.872	1.732	-9.656
1483	CZ	PHE 117	В	36.672	2.294	-B.412
1484	N	ASP 118	В	32.811		-10.426
1485	CA	ASP 118	В	32.630	-5.124	-9.935

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## Fig. 15.34

	tom	Residue	Chain	X	Y_	<b>z</b>
1486	С	ASP 118	В	34.049 -	5.641	-9.774
1487	o	ASP 118	В		4.878	
1488	СВ	ASP 118	В			-10.948
1489	CG	ASP 118	В			-12.291
1490	OD1	ASP 118	B			-13.188
1491	OD2	ASP 118	B			-12.438
1492	N	ASP 119	В		6.943	
1493	CA	ASP 119	В		7.584	-9.692
1494	c	ASP 119	В			-11.043
1495	ŏ	ASP 119	В			-11.918
1496	CB	ASP 119	В		9.105	-9.596
1497	CG	ASP 119	В			-10.579
1498	OD1	ASP 119	В			-11.699
1499	OD2	ASP 119	В	33.370 -1		
1500	N	SER 120	В			-11.190
1501	CA	SER 120	В			-12.116
1502	С	SER 120	В			-11.587
1503	0 -	SER 120	В			-10.396
1504	CB	SER 120	В			-13.505
1505	OG	SER 120	В			-14.048
1506	N	SER 121	В			-12.351
1507	CA	SER 121	В	37.795 -	3.641	-11.746
1508	c	SER 121	В			-12.910
1509	0	SER 121	В			-13.421
1510	СВ	SER 121	В			-11.297
1511	OG	SER 121	В			-10.767
1512	N	ASN 122	В	35.961 -	3.865	-13.437
1513	CA	ASN 122	В	35.251 -	3.163	-14.482
1514	С	ASN 122	В	34.567 -	1.986	-13.818
1515	0	ASN 122	В	33.919 -	2.205	-12.820
1516	CB	ASN 122	В	34.175 -	4.002	-15.124
1517	CG	ASN 122	В	34.720 -	4.905	-16.200
1518	OD1	ASN 122	В	35.807 -	5.455	-16.161
1519	ND2	ASN 122	В		5.095	-17.193
1520	N	VAL 123	В	34.623 -	0.738	-14.229
1521	CA	VAL 123	В	33.707	0.247	-13.704
1522	С	VAL 123	В	32.323	0.092	-14.263
1523	0	VAL 123	В	32.121	0.390	-15.418
1524	CB	VAL 123	В	34.253	1.583	-14.033
1525	CG1	VAL 123	В	33.293	2.705	-13.768
1526	CG2	VAL 123	В	35.490	1.719	-13.184
1527	N	ILE 124	В	31.270 -	0.356	-13.598
1528	CA	ILE 124	В	29.966 -	0.307	-14.250
1529	С	ILE 124	В	28.844	0.618	-13.776
1530	0	ILE 124	В	28.948	1.374	-12.826

Fig. 15.35

	lom	Residue	Chain	<u>x</u>	Y	Z
1531	CB	ILE 124	В	29.448	-1.693	-14.301
1532	CG1	ILE 124	В	29.346	-2.339	-12.994
1533	CG2	ILE 124	В	30.460	-2.470	-15.084
1534	CD1	ILE 124	B	28.941	-3.771	-13.335
1535	N	LEU 125	В.	27.672	0.649	~14.394
1536	CA	LEU 125	В	26.633	1.575	-13.951
1537	С	LEU 125	В	25.470	0.617	-13.807
1538	0	LEU 125	В	24.992	0.201	-14.856
1539	CB	LEU 125	В	26.338	2.598	-15.029
1540	CG	LEU 125	В	25.522	3.838	-14.719
1541	CD1	LEU 125	В	26.188	4.811	-13.745
1542	CD2	LEU 125	В	25.371	4.560	-16.049
1543	N	LYS 126	В	25.012	0.237	-12.583
1544	CA	LYS 126	В	23.788	-0.531	-12.459
1545	С	LYS 126	В	22.674	0.399	-12.077
1546	0	LYS 126	В	22.853	1.502	-11.576
1547	CB	LYS 126	В	23.882	-1.627	-11.411
1548	CG	LYS 126	В	22.987	-2.671	-12.000
1549	CD	LYS 126	В	22.624	-3.980	-11.185
1550	CE	LYS 126	В	21.336		-11.620
1551	NZ	LYS 126	В	20.100	-3.953	-11.514
1552	N	LYS 127	В	21.457		-12.325
1553	CA	LYS 127	В	20.275	0.766	-12.115
1554	С	LYS 127	В	19.538		-11.161
1555	0	LYS 127	В	19.442		-11.349
1556	CB	LYS 127	В	19.357		-13.313
1557	CG	LYS 127	В	18.494		-13.043
1558	CD	LYS 127	В	17.595		-14.203
1559	CE	LYS 127	В	16.759		-14.621
1560	NZ	LYS 127	В	15.623		-15.326
1561	N	TYR 128	В	19.018		-10.119
1562	CA	TYR 128	В	18.374	-0.282	-9.120
1563	C	TYR 128	В	17.039	0.270	-9.294
1564	0	TYR 128	В	16.919	1.484	-9.458
1565	CB	TYR 128	В	18.944	0.051	-7.765
1566	CG	TYR 128	В	20.187	-0.723	-7.613
1567	CD1	TYR 128	В	21.334	-0.090	-7.202
1568	CD2	TYR 128	В	20.189	-2.046	-7.962
1569	CE1	TYR 128	В	22.524	-0.785	-7.189
1570	CE2	TYR 128	В	21.354	-2.752	-7.957
1571	CZ	TYR 128	В	22.522	-2.118	-7.590
1572	OH	TYR 128	В	23.726	-2.834	-7.651
1573	N	ARG 129	В	15.995	-0.539	-9.273
1574	CA	ARG 129	В	14.738	0.093	-9.471
1575	С	ARG 129	В	14.148	0.079	-8.127

Fig. 15.36

A	tom	Residuc	Chain	x	Y	<u>z</u>
1576	0	ARG 129	В	. 14.539	-0.696	-7.272
1577	CB	ARG 129	В	13.872		-10.389
1578	CG	ARG 129	В	14.448		-11.194
1579	CD	ARG 129	В	13.392		-12.257
1580	NE	ARG 129	В	12.867	-1.068	-12.880
1581	CZ	ARG 129	В	13.523	-0.465	-13.872
1582	NH1	ARG 129	В	14.600	-1.048	-14.514
1583	NH2	ARG 129	В	13.110	0.783	-14.224
1584	N	ASN 130	В	13.187	0.959	-7.931
1585	CA	ASN 130	В	12.337	0.930	-6.773
1586	С	ASN 130	В	13.101	1.191	-5.580
1587	0	ASN 130	В	12.956	0.581	-4.555
1588	CB	<b>ASN 130</b>	В	11.644	-0.406	-6.596
1589	CG	ASN 130	В	10.513	-0.422	-7.589
1590	OD1	ASN 130	В	9.991	0.615	-7.968
1591	ND2	ASN 130	В	10.106	-1.571	-8.113
1592	N	MET 131	В	13.978	2.163	-5.682
1593	CA	MET 131	В	14.779	2.497	-4.552
1594	С	MET 131	В	14.403	3.805	-3.898
1595	0	MET 131	В	14.888	4.135	-2.829
1596	CB	MET 131	В	16.193	2.489	-5.056
1597	CG	MET 131	В	16.596	1.063	-5.233
1598	SD	MET 131	В	17.774	0.746	-3.920
1599	CE	MET 131	В	17.131	-0.880	-3.651
1600	N	VAL 132	B	13.540	4.602	-4.492
1601	CA	VAL 132	В	13.272	5.952	-4.016
1602	C	VAL 132	В	11.778	6.099	-3.895
1603	0	VAL 132	В	11.075	5.709	-4.791
1604	CB	VAL 132	В	13.804	6.928	-5.035
1605	CG1	VAL 132	В	13.726	8.358	-4.584
1606	CG2	VAL 132	В	15.252	6.549	-5.285
1607	N	VAL 133	В	11.136	6.617	-2.884
1608	CA	VAL 133	В	9.714	6.827	-2.893
1609	С	VAL 133	В	9.534	8.146	-3.591
1610	0	VAL 133	В	10.322	9.039	-3.333
1611	CB	VAL 133	В	9.257	6.869	-1.443
1612	CG1	VAL 133	В	8.030	7.727	-1.166
1613	CG2	VAL 133	В	9.043	5.425	-1.098
1614	N	ARG 134	В	8.526	8.283	-4.462
1615	CA	ARG 134	В	8.074	9.561	-4.979
1616	С	ARG 134	В	6.805	9.998	-4.253
1617	0	ARG 134	В	6.613	11.145	-3.844
1618	CB	ARG 134	В	7.767	9.438	-6.471
1619	CG	ARG 134	В	8.655	10.170	-7.503
1620	CD	ARG 134	В	7.931	10.074	-8.884

Fig. 15.37

Aı	om	Residue	Chain_	X	Y	Z
					n	
1621	NE	ARG 134	В	8.585	10.702	-10.068
1622	CZ	ARG 134	В	9.074	11.973	-10.169
1623	NH1	ARG 134	В	9.514	12.419	-11.395
1624	NH2	ARG 134	В	9.202	12.813	-9.089
1625	N	ALA 135	В	5.839	9.108	-4.056
1626	CA	ALA 135	В	4.734	9.536	-3.247
1627	C	ALA 135	В	4.163	8.375	-2.497
1628	0	ALA 135	В	4.422	7.228	-2.831
1629	CB	ALA 135	В	3.713	10.178	-4.165
1630	N	CYS 136	В	3.370	8.691	-1.459
1631	CA	CYS 136	В	2.737	7.740	-0.529
1632	С	CYS 136	В	1.234	7.935	-0.411
1633	0	CYS 136	В	0.778	9.062	-0.565
1634	СВ	CYS 136	В	3.124	7.893	0.896
1635	SG	CYS 136	B	4.770	8.538	1.134
1636	N	<b>GLY 137</b>	В	0.452	6.910	-0.132
1637	CA	<b>GLY 137</b>	В	-0.882	7.202	0.246
1638	С	<b>GLY 137</b>	В	-1.562	5.941	0.556
1639	0	<b>GLY 137</b>	В	-0.925	4.911	0.422
1640	N	CYS 138	В	-2.836	5.929	0.957
1641	CA	CYS 138	В	-3.377	4.818	1.746
1642	С	CYS 138	В	-3.790	3.683	0.897
1643	0	CYS 138	В	-4.502	4.036	-0.016
1644	CB	CYS 138	В	-4.575	5.261	2.510
1645	SG	CYS 138	B	-4.204	6.910	3.144
1646	N	HIS 139	В	-3.491	2.408	1.017
1647	CA	HIS 139	В	-4.209	1.485	0.173
1648	С	BIS 139	В	-4.475	0.369	1.139
1649	0	HIS 139	В	-3.753	0.227	2.148
1650	CB	HIS 139	В	-3.407	0.969	-1.034
1651	CG	HIS 139	В	-2.881	2.085	-1.888
1652	ND1	HIS 139	В	-2.348	3.285	-1.424
1653	CD2	HIS 139	В	-3.320	2.198	-3.200
1654	CE1	HIS 139	В	-2.505	4.047	-2.523
1655	NE2	HIS 139	В	-3.077	3.480	-3.620
1656	OT	HIS 139	В	-5.079	-0.465	0.767

Fig. 16\1

A	tom	Resi	duc	Chain	x	Y	Z	<u> </u>
				_				
1	CB	GLN	36	A	34.688	54.268	11.979	0.80
2	CG	GLN	36	A	34.454	55.622	11.327	0.98
3	CD	GLN	36	A	35.339	55.912	10.142	1.07
4	OE1	GLN	36	A	36.525	55.589	10.061	0.98
5	NE2	GLN	36	A	34.724	56.519	9.123	1.05
6	С	GLN	36	A	33.292	52.214	11.583	0.62
7	0	GLN	36	A	33.249	52.268	10.349	0.67
8	N	GLN	36	A	33.249	53.240	13.781	0.82
9	CA	GLN	36	A	33.430	53.511	12.360	0.72
10	N	ALA	37	λ	33.283	51.110	12.314	0.65
11	CA	ALA	37	<b>A</b>	33.203	49.784	11.644	0.64
12	CB	ALA	37	A	33.628	48.665	12.576	0.58
13	С	ALA	37	A	31.758	49.525	11.201	0.62
14	0	ALA	37	A	30.825	49.957	11.868	0.67
15	N	CYS	38	A	31.645	48.593	10.285	0.64
16	CA	CYS	38	A	30.373	48.172	9.714	0.53
17	C	CYS	38	A	29.433	47.699	10.797	0.63
18	0	CYS	38	A	29.746	46.761	11.527	0.66
19	CB	CYS	38	A	30.559	47.212	8.582	0.58
20	SG	CYS	38	A	29.056	46.598	7.828	0.63
21	N	LYS	39	A	28.382	48.487	11.010	0.63
22	CA	LYS	39	A	27.261	48.136	11.860	0.50
23	CB	LYS	39	A	27.365	48.642	13.296	0.56
24	CG	LYS	39	A	28.183	49.898	13.549	0.69
25	CD	LYS	39	A	27.844	50.564	14.870	0.62
26	CE	LYS	39	A	26.701	49.872	15.592	0.92
27	NZ	LYS	39	A	26.052	50.747	16.607	0.92
28	С	LYS	39	A	25.926	48.637	11.262	0.50
29	0	LYS	39	A	25.872	49.334	10.242	0.49
30	N	LYS	40	A	24.890	48.324	11.988	0.59
31	CA	LYS	40	A	23.483	48.556	11.659	0.64
32	CB	LYS	40	A	22.606	47.362	12.132	0.61
33	CG	LYS	40	A	21.167	47.575	11.621	0.60
34	CD	LYS	40	A	20.283	46.452	12.085	0.48
35	CE	LYS	40	A	19.989	46.487	13.565	0.60
36	NZ	LYS	40	A	19.214	45.258	13.873	0.73
37	С	LYS	40	A	23.019	49.744	12.504	0.63
38	0	LYS	40	A	23.463	49.862	13.659	0.61
39	N	HIS	41	A	22.524	50.763	11.826	0.65
40	CA	HIS	41	A	22.150	51.977	12.590	0.58
41	CB	HIS	41	A	22.881	53.195	12.011	0.57
42	CG	HIS	41	A	24.359	52.907	11.899	0.60
43	CD2	HIS	41	A	25.094	52.514	10.838	0.63
44	ND1	HIS	41	A	25.270	53.346	12.835	0.61
45	CE1	HIS	41	A	26.482	53.207	12.338	0.63

Fig. 16\2

A	tom	Resi	due	Chain	x	Υ	z	δ
	V=0	<b></b>		•	06 411	50 515		
46	NE2	HIS	41	A	26.411	52.515	11.217	0.67
47	C	HIS	41	A	20.645	52.146	12.505	0.58
48	0	HIS	41	Α .	19.994	51.655	11.564	0.60
49	N	GLU	42	· A	20.173	53.007	13.373	0.63
50	CA	GLU	42	A	18.747	53.323	13.460	0.65
51	CB	GLU	42	A	18.389	53.683	14.894	0.67
52	CG	GLU	42	A	17.998	52.443	15.724	0.61
53	CD	GLU	42	A	17.770	52.760	17.171	0.80
54	OE1	GLU	42 42	A	18.180	52.068	18.082	0.89
55	OE2	GLU		A	17.221	53.875	17.309	0.92
56	C	GLU	42	A A	18.296	54.397	12.507	0.63
57	0	GLU	42		19.083	55.313 54.191	12.261	0.68
58	И	LEU	43	A	17.163		11.858	0.57
59	CA	LEU	43	A	16.444 16.827	55.149 54.887	11.012 9.561	0.57 0.56
60	CB	LEU	43 43	A	16.795	55.850	8.455	0.65
61	CG	LEU		A				
62	CD1	LEU	43	A	16.623	55.408	7.043	0.59
63	CD2	LEU	43	A	16.638	57.296	8.715	0.64
64	C	LEU	43	A	14.940	54.753	11.210	0.63
65	0	LEU	43	A	14.568	53.682	10.710	0.62
66	N	TYR	44	A	14.119	55.703	11.580	0.58
67	CA	TYR	44	A	12.646	55.504	11.590	0.54
68	CB	TYR	44	A	12.117	56.253	12.845	0.52
69	CG	TYR	44	A	10.648	55.954	13.022	0.63
70	CD1	TYR	44	A	9.663	56.853	12.620	0.50
71	CE1	TYR	44	A	8.318	56.490	12.742	0.58
72	CD2	TYR	44	A	10.267	54.766	13.634	0.52
73	CE2	TYR	44	A	8.957	54.379	13.694	0.56
74	CZ	TYR	44	A	7.976	55.253	13.235	0.69
75	OH	TYR	44	A	6.668	54.894	13.434	0.81
76	C	TYR	44	A	12.114	56.295	10.395	0.51
77	0	TYR	44	A	12.538	57.447	10.316	0.63
78	N	VAL	45	A	11.336	55.713	9.534	0.53
79	CA	VAL	45	A	10.722	56.389	8.368	0.53
80	CB	VAL	45	A	10.910	55.465	7.148	0.56
81	CG1	VAL	45	A	10.355	56.000	5.844	0.47
82	CG2	VAL	45	A	12.346	54.969	7.035	0.52
83	С	VAL	45	A	9.220	56.590	8.673	0.55
84	0	VAL	45	λ	8.585	55.655	9.193	0.53
85	N	SER	46	A	8.851	57.845	8.765	0.59
86	CA	SER	46	A	7.518	58.414	8.858	0.52
87	CB	SER	46	A	7.518	59.907	9.287	0.46
88	QG	SER	46	A	6.397	60.090	10.135	0.69
89	С	SER	46	A	6.883	58.405	7.482	0.52
90	0	SER	46	A	7.484	58.947	6.553	0.56

Fig. 16\3

A	tom	Resi	due	Chain	x	Y	Z	<u> </u>
91	N	PHE	47	A	5.713	57.819	7.380	0.59
92	CA	PHE	47	A	5.009	57.837	6.069	0.57
93	CB	PHE	47	A .	3.844	56.851	6.088	0.53
94	CG	PHE	47	A	4.154	55.395	6.174	0.55
95	CD1	PHE	47	A	5.279	54.859	5.517	0.45
96	CD2	PHE	47	A	3.287	54.509	6.818	0.54
97	CE1	PHE	47	A	5.580	53.514	5.633	0.71
98	CE2	PHE	47	A	3.618	53.164	7.010	0.56
99	CZ	PHE	47	A	4.739	52.644	6.313	0.60
100	С	PHE	47	A	4.737	59.281	5.648	0.49
101	0	PHE	47	A	4.748	59.578	4.426	0.61
102	N	ARG	48	A	4.422	60.187	6.541	0.51
103	CA	ARG	48	A	4.356	61.641	6.314	0.57
104	CB	ARG	48	A	4.183	62.463	7.564	0.49
105 106	CG CD	ARG ARG	48 48	A A	3.277 2.096	62.040 62.917	8.662 8.844	0.74
105	NE	ARG	48	A	2.148	63.816	9.964	0.83 0.71
107	CZ	ARG	48	A	1.255	64.003	10.923	0.66
109	NH1	ARG	48	A	-0.043	63.685	10.923	0.68
110	NH2	ARG	48	A	1.739	64.330	12.123	0.61
111	C	ARG	48	A	5.516	62.216	5.517	0.64
112	0	ARG	48	A	5.342	62.761	4.397	0.69
113	n	ASP	49	A A	6.737	61.953	5.977	0.60
114	CA	ASP	49	A	7.962	62.298	5.274	0.57
115	CB	ASP	49	A	9.205	61.850	6.047	0.68
116	CG	ASP	49	A	9.188	62.342	7.481	0.80
117	OD1	ASP	49	A	10.062	61.939	8.272	0.89
118	OD2	ASP	49	A	8.235	63.099	7.801	0.83
119	c	ASP	49	A	7.998	61.913	3.814	0.66
120	ō	ASP	49	A	8.811	62.490	3.062	0.72
121	N	LEU	50	A	7.456	60.746	3.465	0.70
122	CA	LEU	50	A	7.487	60.243	2.086	0.60
123	CB	LEU	50	Α	7.446	58.703	2.187	0.62
124	CG	LEU	50	A `	8.580	58.098	3.022	0.69
125	CD1	LEU	50	A	8.786	56.662	2.544	0.66
126	CD2	LEU	50	A	9.836	58.911	2.667	0.66
127	С	LEU	50	A	6.275	60.748	1.275	0.62
128	0	LEU	50	A	6.080	60.203	0.167	0.69
129	N	GLY	51	A	5.315	61.265	2.005	0.62
130	CA	GLY	51	A	3.950	61.540	1.563	0.59
131	C	GLY	51	A	3.215	60.231	1.303	0.68
132	0	GLY	51	A	2.960	59.916	0.130	0.73
133	N	TRP	52	A	3.076	59.385	2.325	0.60
134	CA	TRP	52	A	2.385	58.104	2.063	0.63
135	CB	TRP	52	A	3.281	56.916	2.022	0.76

Fig. 16\4

Aı	om	Resi	duc	Chain	X	Y	Z	<u>δ</u>
			<b>5</b> 2	•	4.282	56.698	0.951	0.80
136	CG	TRP	52 52	A A	5.364	55.744	1.004	0.74
137	CD2	TRP	52 52	A	5.991	55.750	-0.263	0.92
138	CE2	TRP	52 52	λ	5.811	54.835	1.963	1.01
139	CE3	TRP	52 52	Α	4.256	57.163	-0.333	0.82
140	CD1	TRP		λ	5.315	56.649	-1.050	0.73
141	NE1	TRP	52	λ	7.038	54.881	-0.576	0.75
142	CZ2	TRP	52		6.886	54.012	1.684	0.68
143	CZ3	TRP	52	Α	7.495	54.041	0.428	0.80
144	CH2	TRP	52	A	1.218	57.994	3.046	0.72
145	С	TRP	52	A	0.425	57.045	2.965	0.72
146	0	TRP	52	λ		59.076	3.769	0.64
147	N	GLN	53	A	1.009 0.000	59.125	4.822	0.74
148	CA	GLN	53	λ		60.290	5.798	0.68
149	CB	GLN	53	A	0.235		5.138	0.56
150	CG	GLN	53	A	0.108	61.647 62.152	4.268	0.78
151	CD	GLN	53	A	1.210	61.501	3.363	0.75
152	OE1	GLN	53	A	1.734			0.80
153	NE2	GLN	53	A	1.476	63.465	4.447	0.50
154	С	GLN	53	A	-1.419	59.100	4.280	
155	0	GLN	53	A	-2.376	59.017	5.080	0.83
156	N	ASP	54	A	-1.613	59.500	3.040	0.77
157	CA	ASP	54	A	-2.945	59.676	2.432	0.76
158	CB	ASP	54	A	-2.747	60.190	1.004	0.92
159	CG	ASP	54	A	-2.418	61.667	0.920	0.92
160	OD1	ASP	54	A	-2.672	62.439	1.863	0.96
161	OD2	ASP	54	A	-2.074	62.062	-0.217	0.75
162	С	ASP	54	A	-3.808	58.408	2.491	0.80
163	0	ASP	54	A	-4.988	58.426	2.898	0.82
164	И	TRP	55	A	-3.265	57.334	1.953	0.72
165	CA	TRP	55	A	-3.905	56.052	1.723	0.78
166	CB	TRP	55	A	-3.527	55.614	0.293	0.69
167	CG	TRP	55	A	-3.900	56.679	-0.691	0.80
168	CD2	TRP	55	A	-4.955	57.641	-0.465	0.76
169	CE2	TRP	55	A	-4.935	58.522	-1.569	0.85
170	CE3	TRP	55	A	-6.064	57.611	0.374	0.87
171	CD1	TRP	55	A	-3.230	57.113	-1.796	0.74
172	NE1	TRP	55	A	-3.835	58.229	-2.337	0.76
173	CZ2	TRP	55	A	-5.913	59.493	-1.745	0.78
174	CZ3	TRP	55	A	-7.074	58.520	0.151	0.88
175	CH2	TRP	55	A	-7.012	59.424	-0.905	0.82
176	С	TRP	55	, A	-3.640	54.977	2.770	0.84
177	0	TRP	55	A	-4.364	53.952	2.758	0.77
178	N	ILE	56	A	-2.653	55.170	3.642	0.83
179	CA	ILE	56	A	-2.272	54.240	4.693	0.78
180	CB	ILE	56	A	-0.750	54.279	5.076	0.77

Fig. 16\5

A	Atom		duc	Chain	<u> </u>	Y	Z	δ
				_				
181	CG2	ILE	56	A	-0.539	54.069	6.606	0.71
182	CG1	ILE	56	A	-0.007	53.147	4.316	0.82
183	CD	ILE	56	A	1.044	53.678	3.298	0.90
184	С	ILE	56	A	-3.122	54.432	5.944	0.74
185	0	ILE	56	A	-3.277	55.575	6.388	0.80
186	N	ILE	57	A	-3.481	53.314	6.567	0.68
187	CA	ILE	57	A	-4.235	53.305	7.822	0.64
188	CB	ILE	57	λ	-5.462	52.319	7.725	0.65
189	CG2	ILE	57	A	-6.003	51.940	9.109	0.55
190	CG1	ILE	57	A	-6.532	52.829	6.736	0.64
191	CD	ILE	57	A	-7.736	51.860	6.540	0.62
192	С	ILE	57	A	-3.350	52.989	9.020	0.60
193	0	ILE	57	A	-3.634	53.435	10.145	0.62
194	N	ALA	58	A	-2.432	52.035	8.834	0.58
195	CA	ALA	58	A	-1.568	51.588	9.967	0.49
196	CB	ALA	58	A	-2.381	50.553	10.789	0.55
197	С	ALA	58	A	-0.372	50.826	9.376	0.46
198	0	ALA	58	A	-0.718	50.311	8.302	0.49
199	и	PRO	59	A	0.849	51.149	9.744	0.53
200	· CD	PRO	59	A	2.056	50.542	9.192	0.46
201	CA	PRO	59	A	1.271	52.004	10.870	0.45
202	CB	PRO	59	A	2.504	51.340	11.429	0.47
203	CG	PRO	59	A	3.139	50.681	10.217	0.43
204	С	PRO	59	A	1.496	53.434	10.384	0.53
205	0	PRO	59	A	1.157	53.822	9.239	0.55
206	N	GLU	60	A	2.009	54.297	11.255	0.49
207	CA	GLU	60	A	2.315	55.678	10.776	0.58
208	CB	GLU	60	A	1.953	56.668	11.906	0.64
209	CG	GLU	60	A	0.536	57.221	11.886	0.77
210	CD	GLU	60	A	-0.327	57.374	13.082	0.79
211	OE1	GLU	60	A	-1.419	56.801	13.170	0.84
212	OE2	GLU	60	A	0.122	58.112	13.997	0.61
213	С	GLU	60	A	3.778	55.806	10.374	0.53
214	0	GLU	60	A	4.180	56.692	9.605	0.52
215	N	GLY	61	A	4.521	54.712	10.496	0.52
216	CA	GLY	61	A	5.866	54.512	9.965	0.54
217	С	GLY	61	A	6.549	53.304	10.585	0.58
218	0	GLY	61	A	5.879	52.548	11.298	0.55
219	N	TYR	62	A	7.807	53.047	10.231	0.55
220	CA	TYR	62	A	8.462	51.838	10.845	0.54
221	CB	TYR	62	A	8.178	50.656	9.853	0.38
222	CG	TYR	62	A	8.980	50.842	8.558	0.47
223	CD1	TYR	62	A	8.538	51.787	7.649	0.47
224	CE1	TYR	62	λ	9.247	52.151	6.499	0.51
225	CD2	TYR	62	A	10.091	50.092	8.201	0.45

Fig. 16\6

At	om	Resi	duc	Chain	x	Y		δ
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226	CE2	TYR	62	A:	10.662	50.225	6.912	0.35
227	CZ	TYR	62	A	10.294	51.305	6.115	0.51
228	OH	TYR	62	A	10.785	51.429	4.839	0.65
229	С	TYR	62	A	9.960	52.092	10.973	0.59
230	0	TYR	62	A	10.580	52.978	10.330	0.55
231	N	ALA	63	A	10.590	51.236	11.731	0.54
232	CA	ALA	63	A	12.067	51.311	11.984	0.49
233	CB	ALA	63	A	12.243	50.507	13.282	0.58
234	С	ALA	63	A	12.734	50.563	10.845	0.51
235	0	ALA	63	A	12.508	49.358	10.665	0.58
236	N	ALA	64	A	13.289	51.297	9.895	0.54
237	CA	ALA	64	A	13.878	50.657	8.735	0.53
238	CB	ALA	64	A	13.779	51.617	7.550	0.53
239	С	ALA	64	A	15.319	50.155	8.972	0.52
240	0	ALA	64	A	15.797	49.399	8.091	0.54
241	N	TYR	65	A	16.092	50.933	9.671	0.51
242	CA	TYR	65	A	17.512	50.804	9.978	0.54
243	CB	TYR	∙65	A	17.944	49.390	10.438	0.47
244	CG	TYR	65	A	17.297	49.063	11.777	0.47
245	CD1	TYR	65	A	17.763	49.564	12.974	0.44
246	CE1	TYR	65	A	17.164	49.219	14.190	0.47
247	CD2	TYR	65	A	15.978	48.600	11.756	0.61
248	CE2	TYR	65	A	15.250	48.484	12.922	0.44
249	CZ	TYR	65	A	15.849	48.733	14.139	0.67
250	OB	TYR	65	A	15.077	48.514	15.253	0.70
251	С	TYR	65	A	18.332	51.188	8.752	0.51
252	0	TYR	65	A	17.714	51.205	7.664	0.49
253	N	TYR	66	A	19.638	51.001	8.928	0.52
254	CA	TYR	66	A	20.521	51.078	7.729	0.50
255	CB	TYR	66	A	20.564	52.484	7.126	0.60
256	CG	TYR	66	A	21.276	53.507	7.983	0.59
257	CD1	TYR	66	A	20.563	54.255	8.917	0.59
258	CE1	TYR	66	A	21.155	55.260	9.659	0.53
259	CD2	TYR	66	A	22.592	53.893	7.721	0.53
260	CE2	TYR	66	λ	23.149	55.001	8.335	0.47
261	CZ	TYR	66	λ	22.511	55.532	9.448	0.58
262	OH	TYR	66	A	23.095	56.600	10.066	0.70
263	C	TYR	66	A	21.939	50.673	8.166	0.51
264	0	TYR	66	A	22.223	50.678	9.351	0.52 0.58
265	N	CYS	67	A	22.764	50.444	7.173	0.58
266	CA	CYS	67	A	24.127	49.897	7.372 7.066	0.57
267	С	CYS	67	A	25.134	50.997		0.32
268	0	CYS	67	A	25.035	51.589	5.968	0.48
269	CB	CXS	67	A	24.270	48.712	6.413	0.54
270	SG	CYS	67	A	23.285	47.244	6.831	0.34

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PCT/US97/01071

Fig. 16\7

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A	tom	Resi	duc	Chain	x_	Y	Z	<u> 8</u>
271	N	GLU	68	A	26.124	51.140	7.930	0.56
272	CA	GLU	68	A	27.251	52.049	7.617	0.47
273	CB	GLD	68	A	26.880	53.495	7.993	0.68
274	CG	GLU	68	A	<b>27.57</b> 5	54.628	7.264	0.53
275	CD	GLU	68	A	27.058	56.019	7.510	0.67
276	OE1	GLU	68	A	27.235	56.925	6.720	0.64
277	OE2	GLU	68	A	26.696	56.240	8.687	0.63
278	С	GLU	68	A	28.467	51.725	8.485	0.43
279	0	GLU	68	A	28.311	51.441	9.663	0.42
280	N	GLY	69	A	29.646	52.013	7.922	0.48
281	CA	GLY	69	A	30.920	51.860	8.669	0.44
282	С	GLY	69	A	31.899	51.218	7.670	0.51
283	0	GLY	69	A	31.525	50.790	6.563	0.53
284	N	GLU	70	A	33.165	51.150	8.048	0.57
285	CA	GLU	70	A	34.166	50.669	7.116	0.63
286	CB	GLU	70	A	35.428	51.440	6.924	0.74
287	CG	GLU	70	A	36.419	51.567	8.062	0.77
288	CD	GLU	70	A	37.808	51.898	7.554	1.02
289	OE1	GLU	70	A	38.022	52.115	6.372	0.84
290	OE2	GLU	70	A	38.665	51.734	8.441	0.95
291	С	GLU	70	A	34.384	49.172	7.133	0.44
292	0	GLU	70	A	34.224	48.564	8.186	0.42
293	N	CYS	71	A	34.793	48.707	5.962	0.50
294	CA	CYS	71	A	35.136	47.281	5.805	0.62
295	C	CYS	71	A	36.652	47.128	5.637	0.54
296	0	CYS	71	A	37.173	47.188	4.547	0.51
297	CB	CYS	71	A	34.283	46.609	4.771	0.62
298	SG	CYS	71	A	32.525	46.435	5.196	0.57
299	N	ALA	72	A	37.345	47.031	6.745	0.50
300	CA	ALA	72	A	38.781	47.033	6.862	0.61
301	CB	ALA	72	A	39.243	48.414	7.425	0.61
302	С	ALA	72	A	39.174	45.969	7.903	0.62
303	0	ALA	72	A	38.444	45.747	8.880	0.59
304	N	PHE	73	A	40.472	45.646	7.785	0.59
305	CA	PHE	73	A	41.098	44.789	8.805	0.46
306	CB	PHE	73	A	42.425	44.197	8.472	0.54
307	CG	PHE	73	A	42.433	43.361	7.198	0.45
308	CD1	PHE	73	A	41.659	42.210	7.129	0.43
309	CD2	PHE	73	A	43.175	43.752	6.117	0.45
310	CE1	PHE	73	A	41.516	41.496	5.935	0.34
311	CE2	PHE	73	A	43.202	42.987	4.959	0.37
312	ÇZ	PHE	73	A	42.340	41.869	4.895	0.33
313	C,	PHE	73	A	40.895	45.531	10.083	0.52
314	0	PHE	73	A	41.336	46.672	9.925	0.54
315	N	PRO	74	A	40.737	44.872	11.197	0.57

Fig. 16\8

	tom	Resi	duc	Chain	<u> </u>	Y	Z	δ
316	CD	PRO	74	A	40.751	45.550	12.495	0.57
317	CA	PRO	74	A	40.171	43.560	11.370	0.56
318	CB	PRO	74	Α .	40.235	43.204	12.832	0.52
319	CG	PRO	74	A	40.362	44.522	13.519	0.54
320	C	PRO	74	A	38.776	43.425	10.805	0.66
321	ō	PRO	74	A	37.919	44.259	11.112	0.74
322	N	LEU	75	A	38.607	42.334	10.090	0.58
323	CA	LEU	75	A	37.302	41.889	9.627	0.61
324	CB	LEU	75	A	37.400	41.397	8.209	0.63
325	CG	LEU	75	A	37.038	42.169	7.006	0.60
326	CD1	LEU	75	A	36.742	43.625	7.025	0.57
327	CD2	LEU	75	A	37.596	41.719	5.703	0.49
328	С	LEU	75	A	36.805	40.946	10.726	0.68
329	0	LEU	75	A	36.983	39.722	10.655	0.72
330	N	ASN	76	A	36.381	41.589	11.802	0.68
331	CA	ASN	76	A	35.736	40.919	12.939	0.66
332	CB	ASN	76	Α	35.320	41.901	14.028	0.62
333	CG	ASN	76	A	36.481	42.611	14.697	0.83
334	OD1	ASN	76	A	37.623	42.110	14.677	0.97
335	ND2	ASN	76	A	36.246	43.740	15.370	0.81
336	C	ASN	76	A	34.658	39.972	12.449	0.67
337	0	ASN	76	A	34.194	39.990	11.281	0.73
338	N	SER	77	A	34.415	38.958	13.260	0.73
339	CA	SER	77	A	33.432	37.909	12.886	0.74
340	СВ	SER	77	A	33.434	36.747	13.894	0.54
341	OG	SER	77	A	33.117	35.573	13.128	0.96
342	С	SER	77	A	32.033	38.539	12.767	0.62
343	0	SER	77	A	31.287	38.245	11.835	0.66
344	N	TYR	78	A	31.705	39.321	13.767	0.63
345	CA	TYR	78	А	30.579	40.220	13.850	0.75
346	CB	TYR	78	А	30.615	41.057	15.137	0.63
347	CG	TYR	78	A	30.922	42.527	14.976	0.91
348	CD1	TYR	78	A	32.115	43.118	15.447	0.81
349	CE1	TYR	78	A	32.383	44.471	15.228	0.88
350	CD2	TYR	78	A	29.929	43.382	14.494	0.98
351	CE2	TYR	78	A	30.178	44.734	14.265	0.99
352	CZ	TYR	78	A	31.402	45.277	14.640	0.94
353	OH	TYR	78	' A	31.541	46.629	14.483	0.97
354	C	TYR	78	A	30.340	41.045	12.592	0.78
355	o	TYR	78	A	29.244	41.627	12.487	0.79
356	N	MET	79	A	31.322	41.194	11.719	0.74
357	CA	MET	79	A	31.228	41.956	10.475	0.62
358	CB	MET	79	A	32.506	42.736	10.191	0.68
359	CG	MET	79	A	32.600	43.844	11.207	0.51
360	SD	MET	79	A	34.243	44.508	11.149	0.67

Fig. 16\9

A	tom	Resi	due	Chain	X	<u>Y</u>	Z	ð
361	CE	MET	79	A	34.390	45.358	9.625	0.53
362	С	MET	79	A	30.807	41.104	9.309	0.58
363	0	MET	79	A	30.601	41.571	8.186	0.61
364	N	ASN	80	A	30.496	39.834	9.575	0.55
365	CA	ASN	80	A	30.017	39.029	8.431	0.54
366	CB	ASN	80	A	28.501	38.847	8.469	0.86
367	CG	ASN	80	A	27.937	38.129	7.244	1.05
368	OD1	ASN	80	A	28.023	38.584	6.079	0.95
369	ND2	ASN	80	A	27.180	37.052	7.518	0.86
370	С	asn	80	A	30.573	39.451	7.091	0.57
371	0	ASN	80	A	29.821	39.382	6.054	0.67
372	N	ALA	81	A	31.852	39.123	6.890	0.60
373	CA	ALA	81	A	32.527	39.331	5.577	0.52
374	CB	ALA	81	A	34.012	39.611	5.894	0.53
375	С	ALA	81	A	32.276	38.077	4.802	0.53
376	0	ALA	81	A	32.376	37.027	5.445	0.60
377	N	THR	82	A	32.057	38.098	3.500	0.54
378	CA	THR	82	A	32.162	36.889	2.680	0.44
379	СВ	THR	82	A	31.426	37.170	1.326	0.45
380	OG1	THR	82	A	32.057	38.409	0.821	0.49
381	CG2	THR	82	A	29.937	37.606	1.601	0.58
382	С	THR	82	A	33.652	36.578	2.443	0.67
383	0	THR	82	A	34.576	37.324	2.852	0.55
384	N	ASN	83	A	33.946	35.396	1.913	0.61
385	CA	ASN	83	A	35.352	35.123	1.529	0.56
386	CB	ASN	83	A	35.532	33.739	0.901	0.53
387	CG	ASN	83	A	35.404	32.602	1.900	0.48
388	OD1	ASN	83	A	35.542	32.820	3.108	0.55
389	ND2	ASN	83	A	34.880	31.483	1.410	0.55
390	С	ASN	83	A	35.756	36.228	0.535	0.55
391	0	ASN	83	A	36.971	36.442	0.428	0.54
392	N	HIS	84	A	35.021	36.349	-0.569	0.55
393	CA	HIS	84	A	35.225	37.440	-1.518	0.41
394	CB	HIS	84	A	34.045	37.721	-2.462	0.58
395	CG	HIS	84	A	34.351	38.672	-3.572	0.40
396	CD2	HIS	84	A	34.775	38.378	-4.809	0.29
397	ND1	HIS	84	A	34.471	40.064	-3.426	0.22
398	CE1	HIS	84	A	34.778	40.549	-4.612	0.18
399	NE2	HIS	84	A	34.912	39.565	-5.513	0.31
400	С	HIS	84	A	35.639	38.754	-0.878	0.47
401	0	HIS	84	A	36.412	39.493	-1.484	0.50
402	N	ALA	85	A	35.014	39.218	0.190	0.52
403	CA	ALA	85	A	35.340	40.514	0.773	0.53
404	CB	ALA	85	A	34.341	40.834	1.868	0.48
405	С	ALA	85	· A	36.756	40.456	1.388	0.57

Fig. 16\10

A	tom	Resi	duc	Chain	x	Y	Z	δ_
44.6	_			•	27 222	40 504		
406	0	ALA	85	Ä	37.222	41.564	1.781	0.55
407	N	ILE	86	A	37.011	39.347	2.095	0.50
408	CA	ILE	86	A	38.305	39.171	2.783	0.52
409	СВ	ILE	86	A	38.390	37.738	3.425	0.57
410	CG2	ILE	86	A	39.877	37.505	3.912	0.42
411	CG1	ILE	86	A	37.435	37.747	4.652	0.40
412	CD	ILE	86	A	37.270	36.306	5.234	0.49
413	С	ILE	86	A	39.432	39.316	1.751	0.45
414	0	ILE	86	A	40.345	40.113	1.894	0.58
415	И	VAL	87	A	39.232	38.678	0.618	0.49
416	CA	VAL	87	A	40.135	38.769	-0.517	0.42
417	CB	VAL	87	A	39.663	37.786	-1.570	0.43
418	CG1	VAL	87	A	40.259	38.002	-2.950	0.39
419	CG2	VAL	87	A	39.952	36.365	-1.097	0.50
420	С	VAL	87	A	40.276	40.212	-0.984	0.52
421	0	VAL	87	A	41.217	40.442	-1.765	0.57
422	И	GLN	88	A	39.086	40.754	-1.301	0.64
423	CA	GLN	88	A	38.956	42.130	-1.773	0.50
424	CB	GLN	88	A	37.685	42.583	-2.372	0.54
425	CG	GLN	88	A	37.381	44.072	-2.355	0.50
426	CD	GLN	88	A	36.114	44.340	-3.126	0.53
427	OE 1	GLN	88	A	35.371	43.417	-3.448	0.63
428	NE2	GLN	88	λ	35.759	45.595	-3.183	0.42
429	С	GLN	88	A	39.679	43.085	-0.B67	0.42
430	0	GLN	88	A	40.292	44.052	-1.314	0.52
431	N	THR	89	A	39.512	42.927	0.420	0.52
432	CA	THR	89	A	40.198	43.702	1.434	0.51
433	CB	THR	89	A	39.557	43.573	2.830	0.44
434	OG1	THR	89	A	38.120	43.824	2.666	0.58
435	CG2	THR	89	A	40.094	44.473	3.906	0.28
436	С	THR	89	A	41.709	43.490	1.437	0.60
437	0	THR	89	A	42.412	44.454	1.772	0.60
438	N	LEU	90	A	42.218	42.388	0.944	0.60
439	CA	LEU	90	· <b>A</b>	43.647	42.052	0.965	0.54
440	CB	LEU	90	A	43.888	40.582	1.267	0.64
441	CG	LEU	90	Α	45.291	40.015	1.078	0.62
442	CD1	LEU	90	A	46.152	40.465	2.272	0.49
443	CD2	LEU	90	A	45.199	38.484	1.213	0.62
444	С	LEU	90	A	44.288	42.626	-0.279	0.47
445	0	LEU	90	A	45.284	43.352	-0.185	0.63
446	N	VAL	91	A	43.579	42.594	-1.376	0.43
447	CA	VAL	91	A	43.901	43.242	-2.618	0.40
448	CB	VAL	91	A	42.933	42.872	-3.726	0.35
449	CG1	VAL	91	A	43.378	43.386	-5.070	0.33
450	CG2	VAL	91	A	42.891	41.354	-3.941	0.47

Fig. 16\11

A	tom	Resi	due	Chain	<u>x</u>	Y	z	<u> </u>
				_	44 170	44 707	0.505	0.40
451	С	VAL	91	A	44.178	44.727	-2.505	0.49
452	0	VAL	91	A	44.650	45.311	-3.477	0.59
453	И	HIS	92	A	43.441	45.341	-1.635	0.51
454	CA	HIS	92	A	43.261	46.741	-1.409	0.56
455	СВ	HIS	92	A	41.870	47.044	-0.770	0.55
456	CG	HIS	92	A	41.747	48.505	-0.469	0.52
457	CD2	HIS	92	A	42.030	49.193	0.668	0.36
458	ND1	HIS	92	A	41.469	49.429	-1.483	0.42
459	CE1	HIS	92	A	41.437	50.616	-0.876	0.49
460	NE2	HIS	92	A	41.799	50.525	0.379	0.43
461	С	HIS	92	A	44.394	47.223	-0.492	0.50
462	0	HIS	92	A	45.068	48.210	-0.766	0.62
463	N	PHE	93	A	44.696	46.435	0.494	0.52
464	CA	PHE	93	A	45.853	46.579	1.335	0.62
465	CB	PHE	93	A	45.848	45.644	2.519	0.52
466	CG	PHE	93	A	47.210	45.573	3.167	0.65
467	CD1	PHE	93	A	48.069	44.545	2.795	0.60
468	CD2	PHE	93	A	47.593	46.500	4.135	0.66
469	CE1	PHE	93	A	49.393	44.536	3.237	0.64
470	CE2	PHE	93	A	48.928	46.509	4.580	0.66
471	CZ	PHE	93	A	49.711	45.369	4.295	0.51
472	С	PHE	93	A	47.144	46.550	0.506	0.66
473	0	PHE	93	A .	47.943	47.489	0.617	0.73
474	N	ILE	94	A	47.331	45.557	-0.315	0.64
475	CA	ILE	94	A	48.423	45.367	-1.252	0.61
476	CB	ILE	94	A	48.402	43.957	-1.897	0.63
477	CG2	ILE	94	A	49.466	43.645	-2.972	0.63
478	CG1	ILE	94	A	48.404	42.831	-0.814	0.51
479	CD	ILE	94	A	48.147	41.476	-1.602	0.63
480	С	ILE	94	A	48.550	46.477	-2.289	0.70
481	0	ILE	94	A	49.659	46.609	-2.824	0.75
482	N	ASN	95	A	47.465	46.873	-2,923	0.74
483	CA	ASN	95	A	47.356	47.921	-3.913	0.64
484	CB	ASN	95	A	47.834	47.543	-5.288	0.70
485	CG	ASN	95	A	48.029	48.754	-6.197	0.86
486	OD1	asn	95	λ	47.797	49.907	-5.782	0.79
487	ND2	ASN	95	λ	48.255	48.499	-7.487	0.87
488	С	asn	95	A	46.014	48.658	-3.867	0.65
489	0	ASN	95	A	45.144	48.489	-4.746	0.65
490	N	PRO	96	A	46.039	49.731	-3.096	0.64
491	CD	PRO	96	A	47.201	50.067	-2.228	0.68
492	CA	PRO	96	A	44.914	50.589	-2.805	0.65
493	CB	PRO	96	A	45.500	51.713	-1.943	0.69
494	CG	PRO	96	A	46.582	50.992	-1.180	0.72
495	C	PRO	96	A	44.106	51.115	-3.961	0.72

Fig. 16\12

521         CD         PRO         100         A         37.460         48.923         -5.352         0.54           522         CA         PRO         100         A         38.083         49.661         -3.125         0.49           523         CB         PRO         100         A         36.659         50.149         -3.411         0.50           524         CG         PRO         100         A         36.231         49.470         -4.666         0.47           525         C         PRO         100         A         38.085         48.608         -2.041         0.41           526         O         PRO         100         A         37.913         47.407         -2.250         0.58           527         N         LYS         101         A         37.739         49.029         -0.836         0.48           528         CA         LYS         101         A         37.549         48.844         1.590         0.58           530         CG         LYS         101         A         38.756         48.257         2.367         0.69           531         CD         LYS         101         A	At	om	Res	idue	Chain	x	Y	<u>z</u>	δ
497 N GLU 97 A 44.662 50.992 -5.140 0.72 498 CA GLU 97 A 44.135 51.677 -6.338 0.70 499 CB GLU 97 A 45.501 52.282 -7.121 0.84 500 CG GLU 97 A 45.685 53.733 -7.069 0.80 501 CD GLU 97 A 45.685 53.733 -7.069 0.80 501 CD GLU 97 A 45.685 53.733 -7.069 0.80 502 OE1 GLU 97 A 45.685 53.733 -7.069 0.80 503 OE2 GLU 97 A 44.477 55.564 -6.044 1.13 503 OE2 GLU 97 A 46.026 54.515 -4.948 1.11 504 C GLU 97 A 43.468 50.625 -7.224 0.72 505 O GLU 97 A 43.686 50.625 -7.224 0.72 505 O GLU 97 A 43.868 50.625 -7.224 0.72 505 O GLU 97 A 43.868 50.625 -7.224 0.72 505 O GLU 97 A 43.868 50.625 -7.500 0.67 506 N THR 98 A 43.863 49.373 -7.015 0.69 507 CA THR 98 A 43.305 48.280 -7.830 0.67 508 CB THR 98 A 44.071 46.915 -7.570 0.65 509 OG1 THR 98 A 44.197 46.102 -8.854 0.78 510 CG2 THR 98 A 41.917 46.8097 -7.527 0.63 511 C THR 98 A 41.817 48.097 -7.527 0.63 513 N VAL 99 A 41.488 48.234 -6.249 0.57 514 CA VAL 99 A 40.083 47.984 -5.846 0.52 515 CB VAL 99 A 39.921 46.444 -5.871 0.56 516 CG1 VAL 99 A 39.921 46.444 -5.871 0.56 517 CG2 VAL 99 A 38.532 45.932 -6.138 0.62 518 C VAL 99 A 39.913 48.669 -4.532 0.52 519 O VAL 99 A 38.532 45.932 -6.138 0.62 520 N PRO 100 A 38.557 49.141 -4.403 0.53 521 CD PRO 100 A 38.083 49.661 -3.125 0.49 522 CA PRO 100 A 38.083 49.661 -3.125 0.49 523 CB PRO 100 A 36.659 50.149 -3.411 0.50 524 CG PRO 100 A 36.659 50.149 -3.411 0.50 525 C PRO 100 A 37.739 49.029 -0.836 0.48 526 O PRO 100 A 38.085 48.608 -2.041 0.41 529 CB LYS 101 A 37.739 49.029 -0.836 0.48 533 NZ LYS 101 A 37.549 48.844 1.590 0.58 533 NZ LYS 101 A 39.523 51.248 4.589 0.76 533 NZ LYS 101 A 39.523 51.248 4.589 0.76 536 N PRO 102 A 36.655 45.655 1.164 0.48 537 CD PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.655 45.60									
497 N GLU 97 A 44.662 50.992 -5.140 0.72 498 CA GLU 97 A 44.135 51.677 -6.338 0.70 499 CB GLU 97 A 45.501 52.282 -7.121 0.84 500 CG GLU 97 A 45.685 53.733 -7.069 0.80 501 CD GLU 97 A 45.685 53.733 -7.069 0.80 501 CD GLU 97 A 45.685 53.733 -7.069 0.80 502 OE1 GLU 97 A 45.685 53.733 -7.069 0.80 503 OE2 GLU 97 A 44.477 55.564 -6.044 1.13 503 OE2 GLU 97 A 46.026 54.515 -4.948 1.11 504 C GLU 97 A 43.468 50.625 -7.224 0.72 505 O GLU 97 A 43.868 50.625 -7.224 0.72 505 O GLU 97 A 43.868 50.625 -7.224 0.72 506 N THR 98 A 43.863 49.373 -7.015 0.69 507 CA THR 98 A 43.305 48.280 -7.830 0.67 508 CB THR 98 A 44.071 46.915 -7.570 0.65 509 OG1 THR 98 A 44.071 46.915 -7.570 0.65 509 OG1 THR 98 A 41.917 46.102 -8.854 0.78 510 CG2 THR 98 A 41.917 46.102 -8.854 0.78 511 C THR 98 A 41.046 47.740 -8.443 0.67 513 N VAL 99 A 41.488 48.234 -6.249 0.57 514 CA VAL 99 A 40.083 47.984 -5.886 0.52 515 CB VAL 99 A 39.921 46.444 -5.871 0.56 516 CG1 VAL 99 A 39.921 46.444 -5.871 0.56 517 CG2 VAL 99 A 39.921 46.444 -5.871 0.56 518 C VAL 99 A 39.921 46.444 -5.871 0.56 520 N PRO 100 A 38.557 49.141 -4.403 0.53 521 CD PRO 100 A 38.557 49.141 -4.403 0.53 522 CA PRO 100 A 38.083 49.661 -3.125 0.49 524 CG PRO 100 A 38.083 49.661 -3.125 0.49 525 C PRO 100 A 38.083 49.661 -3.125 0.49 526 O PRO 100 A 38.083 49.661 -3.125 0.49 527 N LYS 101 A 37.739 49.029 -0.836 0.48 528 CA LYS 101 A 37.739 49.029 -0.836 0.48 529 CB LYS 101 A 37.549 48.844 1.590 0.58 530 CG LYS 101 A 38.756 48.257 2.367 0.65 531 CD LYS 101 A 39.223 51.248 4.589 0.76 533 NZ LYS 101 A 39.523 51.248 4.589 0.76 533 NZ LYS 101 A 39.523 51.248 4.589 0.76 536 N PRO 102 A 36.555 45.605 1.164 0.48 537 CD PRO 102 A 36.555 45.605 1.164 0.48 539 CB PRO 102 A 36.557 49.144 -6.290 0.429 0.55 530 CB PRO 102 A 36.555 45.605 1.164 0.48 531 CD PRO 102 A 36.659 50.149 -3.411 0.69 533 CB PRO 102 A 36.655 45.605 1.164 0.48 535 O LYS 101 A 39.523 51.248 4.589 0.76 536 N PRO 102 A 36.655 45.605 1.164 0.48 537 CD PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.655 45.605 1.164 0.48 539 CB PRO 102 A 36.6	196	0	PRO	96	A	42.930	51.520	-3.812	0.73
498 CA GLU 97 A 44.135 51.677 -6.338 0.70 499 CB GLU 97 A 45.301 52.282 -7.121 0.84 500 CG GLU 97 A 45.685 53.733 -7.069 0.80 501 CD GLU 97 A 45.685 53.733 -7.069 0.80 502 OE1 GLU 97 A 45.275 54.630 -5.952 1.11 502 OE1 GLU 97 A 46.026 54.515 -4.948 1.11 503 OE2 GLU 97 A 46.026 54.515 -4.948 1.11 504 C GLU 97 A 43.468 50.625 -7.224 0.72 505 O GLU 97 A 43.468 50.625 -7.224 0.72 505 O GLU 97 A 43.863 49.373 -7.015 0.69 507 CA THR 98 A 43.863 49.373 -7.015 0.69 508 CB TER 98 A 44.071 46.915 -7.570 0.65 509 OG1 THR 98 A 44.071 46.915 -7.570 0.65 509 OG1 THR 98 A 44.197 46.102 -8.854 0.78 511 C THR 98 A 41.817 48.097 -7.527 0.63 512 O THR 98 A 41.817 48.097 -7.527 0.63 513 N VAL 99 A 41.488 48.234 -6.249 0.57 514 CA VAL 99 A 40.083 47.984 -5.846 0.52 515 CB VAL 99 A 39.921 46.444 -5.871 0.56 516 CG1 VAL 99 A 39.921 46.444 -5.871 0.56 517 CG2 VAL 99 A 39.921 46.444 -5.871 0.56 518 C VAL 99 A 39.921 46.444 -5.871 0.56 520 N PRO 100 A 37.460 48.923 -6.138 0.62 521 CD PRO 100 A 38.659 50.149 -3.411 0.50 522 CA PRO 100 A 38.659 50.149 -3.411 0.50 524 CG PRO 100 A 37.960 48.923 -5.352 0.54 525 C PRO 100 A 37.931 47.407 -2.250 0.58 526 C PRO 100 A 37.931 47.407 -2.250 0.58 527 N LYS 101 A 37.365 48.284 1.590 0.56 533 CG LYS 101 A 39.462 49.214 3.271 0.65 534 C LYS 101 A 39.462 49.214 3.271 0.65 535 C LYS 101 A 39.462 49.214 3.271 0.65 536 N PRO 102 A 35.434 46.638 0.475 0.59 537 CD PRO 102 A 35.530 44.912 1.028 0.43 538 CA PRO 102 A 35.434 46.638 0.475 0.55 539 CB PRO 102 A 35.434 46.638 0.475 0.55 539 CB PRO 102 A 35.434 46.639 0.475 0.55 539 CB PRO 102 A 35.434 46.639 0.475 0.55 539 CB PRO 102 A 36.655 545.605 1.164 0.58 539 CB PRO 102 A 36.657 54.608 0.475 0.57 539 CB PRO 102 A 35.434 46.639 0.475 0.55 539 CB PRO 102 A 36.625 545.605 1.164 0.58 539 CB PRO 102 A 36.625 545.605 1.164 0.58 539 CB PRO 102 A 36.625 545.605 1.164 0.58 539 CB PRO 102 A 36.625 545.605 1.164 0.58 539 CB PRO 102 A 36.625 545.605 1.164 0.58 539 CB PRO 102 A 36.625 545.605 1.164 0.58							50.992	-5.140	0.72
499 CB GLU 97 A 45.301 52.282 -7.121 0.84 500 CG GLU 97 A 45.685 53.733 -7.069 0.80 501 CD GLU 97 A 45.685 53.733 -7.069 0.80 502 GEI GLU 97 A 45.685 53.733 -7.069 0.80 503 OE2 GLU 97 A 44.477 55.564 -6.044 1.13 503 OE2 GLU 97 A 46.026 54.515 -4.948 1.11 504 C GLU 97 A 43.468 50.625 -7.224 0.72 505 O GLU 97 A 42.759 50.951 -8.192 0.74 506 N THR 98 A 43.863 49.373 -7.015 0.69 507 CA THR 98 A 43.863 49.373 -7.015 0.69 509 OG1 THR 98 A 43.305 48.280 -7.830 0.67 509 OG1 THR 98 A 44.071 46.915 -7.570 0.65 509 OG1 THR 98 A 44.197 46.102 -8.854 0.78 510 CG2 THR 98 A 41.817 48.097 -7.527 0.63 511 C THR 98 A 41.817 48.097 -7.527 0.63 512 O THR 98 A 41.046 47.740 -8.443 0.67 513 N VAL 99 A 41.488 48.234 -6.249 0.57 514 CA VAL 99 A 40.083 47.984 -5.866 0.52 515 CB VAL 99 A 39.921 46.444 -5.871 0.56 516 CG1 VAL 99 A 39.921 46.444 -5.871 0.56 518 C VAL 99 A 39.813 48.669 -4.553 0.62 518 C VAL 99 A 39.813 48.669 -4.553 0.62 519 O VAL 99 A 39.813 48.669 -4.552 0.52 510 CD PRO 100 A 38.557 49.141 -4.403 0.52 522 CA PRO 100 A 38.083 49.661 -3.125 0.49 523 CB PRO 100 A 36.639 50.149 -3.411 0.40 524 CG PRO 100 A 36.639 50.149 -3.411 0.40 525 C PRO 100 A 36.639 50.149 -3.411 0.40 526 O PRO 100 A 37.7460 48.923 -5.5352 0.54 527 N LYS 101 A 37.739 49.029 -0.836 0.47 533 NZ LYS 101 A 37.549 48.844 1.590 0.58 534 C LYS 101 A 39.462 49.214 3.271 0.65 535 C LYS 101 A 39.462 49.214 3.271 0.65 536 N PRO 102 A 35.850 47.825 0.048 0.56 537 CD PRO 102 A 36.655 50.149 -3.411 0.65 538 CB PRO 100 A 36.655 50.149 -3.411 0.65 536 N PRO 102 A 35.850 47.825 0.048 0.56 537 CD PRO 102 A 35.850 47.825 0.048 0.56 538 CB PRO 102 A 36.655 50.149 -3.411 0.65 539 CB PRO 102 A 36.655 50.149 -3.411 0.65 536 N PRO 102 A 35.850 47.825 0.048 0.56 537 CD PRO 102 A 36.655 50.149 -3.411 0.65 538 CB PRO 102 A 36.655 50.149 -3.411 0.65 539 CB PRO 102 A 36.655 50.149 -3.411 0.65 539 CB PRO 102 A 36.655 50.149 -3.411 0.65 539 CB PRO 102 A 36.650 47.825 0.048 0.56 539 CB PRO 102 A 36.650 47.825 0.048 0.56 539 CB PRO 102 A 36.650 47.820 0.628 0.59							51.677	-6.338	0.70
500         CG         GLU         97         A         45.685         53.733         -7.069         0.80           501         CD         GLU         97         A         45.275         54.630         -5.952         1.11           502         OE1         GLU         97         A         44.477         55.564         -6.044         1.13           503         OE2         GLU         97         A         46.026         54.515         -4.948         1.11           504         C         GLU         97         A         43.468         50.625         -7.224         0.72           505         O         GLU         97         A         43.468         50.625         -7.224         0.72           506         N         TERR         98         A         43.305         48.280         -7.830         0.67           507         CA         TERR         98         A         44.071         46.915         -7.570         0.65           509         GL         TERR         98         A         44.071         46.915         -7.570         0.67           510         CG2         TERR         98         A						45.301	52.282	-7.121	0.84
501         CD         GLU         97         A         45.275         54.630         -5.952         1.11           502         OE1         GLU         97         A         44.477         55.564         -6.044         1.11           504         C         GLU         97         A         46.026         54.515         -4.948         1.11           505         O         GLU         97         A         42.759         50.951         -8.192         0.74           506         N         TER         98         A         43.863         49.373         -7.015         0.69           507         CA         TER         98         A         43.863         49.373         -7.015         0.69           509         OG1         TER         98         A         44.071         46.915         -7.570         0.65           509         OG1         TER         98         A         44.071         46.915         -7.570         0.65           509         OG1         TER         98         A         41.817         48.097         -7.527         0.63           510         CG2         TER         98         A						45.685	53.733	-7.069	0.80
502         OE1         GLU         97         A         44.477         55.564         -6.044         1.13           503         OE2         GLU         97         A         46.026         54.515         -4.948         1.11           504         C         GLU         97         A         43.468         50.625         -7.224         0.72           505         O         GLU         97         A         42.759         50.951         -8.192         0.74           506         N         THR         98         A         43.863         49.373         -7.015         0.69           507         CA         THR         98         A         44.071         46.915         -7.570         0.65           509         OG1         THR         98         A         44.197         46.102         -8.846         0.78           510         CG2         THR         98         A         41.046         47.740         -8.846         0.78           511         C         THR         98         A         41.046         47.740         -8.443         0.67           512         O         THR         98         A         <						45.275	54.630	-5.952	1.11
503         OE2         GLU         97         A         46.026         54.515         -4.948         1.11           504         C         GLU         97         A         43.468         50.625         -7.224         0.72           505         O         GLU         97         A         42.759         50.951         -8.192         0.74           506         N         THR         98         A         43.863         49.373         -7.015         0.69           507         CA         THR         98         A         44.071         46.915         -7.570         0.65           509         OG1         THR         98         A         44.071         46.102         -8.54         0.78           510         CG2         THR         98         A         41.817         46.097         -7.527         0.63           511         C         THR         98         A         41.817         46.097         -7.527         0.63           512         O         THR         98         A         41.48         48.234         -6.249         0.57           512         O         THR         98         A         4					A	44.477	55.564	-6.044	1.13
504         C         GLU         97         A         43.468         50.625         -7.224         0.72           505         O         GLU         97         A         42.759         50.951         -8.192         0.74           506         N         THR         98         A         43.863         49.373         -7.015         0.65           507         CA         THR         98         A         43.305         48.280         -7.830         0.67           508         CB         THR         98         A         44.071         46.915         -7.570         0.65           509         OG1         THR         98         A         44.197         46.102         -8.854         0.78           510         CG2         THR         98         A         41.817         46.097         -7.527         0.63           511         C         THR         98         A         41.817         46.097         -7.527         0.63           512         O         THR         98         A         41.046         47.740         -8.443         0.67           513         C         VAL         99         A					A	46.026	54.515	-4.948	1.11
505         O         GLU         97         A         42.759         50.951         -8.192         0.74           506         N         THR         98         A         43.863         49.373         -7.015         0.69           507         CA         THR         98         A         43.305         48.280         -7.830         0.67           509         OGI         THR         98         A         44.071         46.915         -7.570         0.65           509         OGI         THR         98         A         44.071         46.102         -8.854         0.78           510         CG2         THR         98         A         41.817         46.102         -8.854         0.78           511         C         THR         98         A         41.817         46.102         -8.854         0.78           511         C         THR         98         A         41.881         49.097         -7.527         0.63           512         O         THR         98         A         41.948         48.234         -6.249         0.57           513         N         VAL         99         A <th< td=""><td></td><td></td><td></td><td></td><td>A</td><td>43.468</td><td>50.625</td><td>-7.224</td><td>0.72</td></th<>					A	43.468	50.625	-7.224	0.72
506         N         THR         98         A         43.863         49.373         -7.015         0.69           507         CA         THR         98         A         43.305         48.280         -7.830         0.67           508         CB         THR         98         A         44.071         46.915         -7.570         0.65           509         OGI         THR         98         A         44.071         46.102         -8.854         0.78           510         CG2         THR         98         A         41.817         48.097         -7.527         0.63           511         C         THR         98         A         41.817         48.097         -7.527         0.63           512         O         THR         98         A         41.046         47.740         -8.443         0.67           513         N         VAL         99         A         41.488         48.234         -6.249         0.57           514         CA         VAL         99         A         40.083         47.984         -5.846         0.52           515         CB         VAL         99         A <t< td=""><td></td><td></td><td></td><td>97</td><td>A</td><td>42.759</td><td>50.951</td><td>-8.192</td><td>0.74</td></t<>				97	A	42.759	50.951	-8.192	0.74
507         CA         THR         98         A         43.305         48.280         -7.830         0.67           508         CB         THR         98         A         44.071         46.915         -7.570         0.65           509         OG1         THR         98         A         44.339         47.345         -7.123         0.77           510         CG2         THR         98         A         44.197         46.102         -8.854         0.78           511         C         THR         98         A         41.817         48.097         -7.527         0.63           512         O         THR         98         A         41.046         47.740         -8.443         0.67           512         O         THR         98         A         41.488         48.234         -6.249         0.57           514         CA         VAL         99         A         40.083         47.984         -5.846         0.52           515         CB         VAL         99         A         39.921         46.444         -5.871         0.56           516         CG1         VAL         99         A					A	43.863	49.373	-7.015	0.69
508         CB         THR         98         A         44.071         46.915         -7.570         0.65           509         OG1         THR         98         A         45.398         47.345         -7.123         0.77           510         CG2         THR         98         A         44.197         46.102         -8.854         0.78           511         C         THR         98         A         41.817         46.097         -7.527         0.63           512         O         THR         98         A         41.046         47.740         -8.443         0.67           513         N         VAL         99         A         41.488         48.234         -6.249         0.57           514         CA         VAL         99         A         40.083         47.984         -5.866         0.52           515         CB         VAL         99         A         40.467         48.868         -4.573         0.44           517         CG2         VAL         99         A         39.813         48.669         -4.532         0.52           519         O         VAL         99         A         <				98	À	43.305	48.280	-7.830	0.67
509         OG1         THR         98         A         45.398         47.345         -7.123         0.77           510         CG2         THR         98         A         44.197         46.102         -8.854         0.78           511         C         THR         98         A         41.817         48.097         -7.527         0.63           512         O         THR         98         A         41.046         47.740         -8.443         0.67           513         N         VAL         99         A         41.488         48.234         -6.249         0.57           514         CA         VAL         99         A         40.083         47.984         -5.846         0.52           515         CB         VAL         99         A         40.467         45.868         -4.573         0.44           517         CG2         VAL         99         A         38.532         26.138         0.62           518         C         VAL         99         A         39.813         48.669         -4.532         0.52           519         O         VAL         99         A         39.813 <t< td=""><td></td><td></td><td></td><td></td><td>A</td><td>44.071</td><td>46.915</td><td>-7.570</td><td>0.65</td></t<>					A	44.071	46.915	-7.570	0.65
510         CG2         THR         98         A         44.197         46.102         -8.854         0.78           511         C         THR         98         A         41.817         48.097         -7.527         0.63           512         O         THR         98         A         41.081         49.097         -7.527         0.63           513         N         VAL         99         A         41.046         47.740         -8.443         0.67           514         CA         VAL         99         A         40.083         47.984         -5.846         0.52           515         CB         VAL         99         A         39.921         46.444         -5.871         0.56           516         CG1         VAL         99         A         39.813         48.669         -4.573         0.44           517         CG2         VAL         99         A         39.813         48.669         -4.532         0.52           519         O         VAL         99         A         40.679         48.871         -3.666         0.53           520         N         PRO         100         A         <				98	A	45.398	47.345	-7.123	0.77
511         C         THR         98         A         41.817         48.097         -7.527         0.63           512         O         THR         98         A         41.046         47.740         -8.443         0.67           513         N         VAL         99         A         41.488         48.234         -6.249         0.57           514         CA         VAL         99         A         40.083         47.984         -5.846         0.52           515         CB         VAL         99         A         39.921         46.444         -5.871         0.56           516         CG1         VAL         99         A         40.467         45.868         -4.573         0.44           517         CG2         VAL         99         A         39.813         48.669         -4.532         0.52           518         C         VAL         99         A         40.679         48.871         -3.666         0.53           518         C         VAL         99         A         40.679         48.871         -3.460         0.52           518         C         VAL         99         A					A	44.197	46.102	-8.854	0.78
512         O         THR         98         A         41.046         47.740         -8.443         0.67           513         N         VAL         99         A         41.488         48.234         -6.249         0.57           514         CA         VAL         99         A         40.083         47.984         -5.846         0.52           515         CB         VAL         99         A         39.921         46.444         -5.871         0.56           516         CG1         VAL         99         A         40.467         45.868         -4.573         0.44           517         CG2         VAL         99         A         38.532         45.932         -6.138         0.62           518         C         VAL         99         A         39.813         48.669         -4.532         0.52           519         O         VAL         99         A         40.679         48.871         -3.666         0.53           520         N         PRO         100         A         38.083         49.661         -3.125         0.54           521         CD         PRO         100         A         <					A	41.817	48.097	- <b>7</b> .527	0.63
513         N         VAL         99         A         41.488         48.234         -6.249         0.57           514         CA         VAL         99         A         40.083         47.984         -5.846         0.52           515         CB         VAL         99         A         39.921         46.444         -5.871         0.56           516         CG1         VAL         99         A         38.532         45.932         -6.138         0.62           518         C         VAL         99         A         39.813         48.669         -4.532         0.52           519         O         VAL         99         A         40.679         48.871         -3.666         0.53           520         N         PRO         100         A         38.557         49.141         -4.403         0.53           521         CD         PRO         100         A         36.659         50.149         -3.411         0.50           522         CA         PRO         100         A         36.659         50.149         -3.411         0.50           523         CB         PRO         100         A						41.046	47.740	-B.443	0.67
514         CA         VAL         99         A         40.083         47.984         -5.846         0.52           515         CB         VAL         99         A         39.921         46.444         -5.871         0.56           516         CGI         VAL         99         A         40.467         45.868         -4.573         0.44           517         CG2         VAL         99         A         38.532         45.932         -6.138         0.62           518         C         VAL         99         A         39.813         48.669         -4.532         0.52           519         O         VAL         99         A         40.679         48.871         -3.666         0.53           520         N         PRO         100         A         38.557         49.141         -4.403         0.53           521         CD         PRO         100         A         37.460         48.923         -5.352         0.54           522         CA         PRO         100         A         36.659         50.149         -3.411         0.50           524         CG         PRO         100         A				99	A	41.488	48.234	-6.249	0.57
515         CB         VAL         99         A         39.921         46.444         -5.871         0.56           516         CG1         VAL         99         A         40.467         45.868         -4.573         0.44           517         CG2         VAL         99         A         38.532         45.932         -6.138         0.62           518         C         VAL         99         A         39.813         48.669         -4.532         0.52           519         O         VAL         99         A         40.679         48.871         -3.666         0.53           520         N         PRO         100         A         38.557         49.141         -4.403         0.53           521         CD         PRO         100         A         37.460         48.923         -5.352         0.54           522         CA         PRO         100         A         36.659         50.149         -3.411         0.50           524         CG         PRO         100         A         36.231         49.470         -4.666         0.47           525         C         PRO         100         A				99	A	40.083	47.984	-5.846	0.52
516         CG1         VAL         99         A         40.467         45.868         -4.573         0.44           517         CG2         VAL         99         A         38.532         45.932         -6.138         0.62           518         C         VAL         99         A         39.813         48.669         -4.532         0.52           519         O         VAL         99         A         40.679         48.871         -3.666         0.53           520         N         PRO         100         A         38.557         49.141         -4.403         0.53           521         CD         PRO         100         A         37.460         48.923         -5.352         0.54           522         CA         PRO         100         A         38.083         49.661         -3.125         0.49           523         CB         PRO         100         A         36.559         50.149         -3.411         0.50           524         CG         PRO         100         A         38.085         48.608         -2.041         0.41           525         C         PRO         100         A					A	39.921	46.444	-5.871	0.56
517         CG2         VAL         99         A         38.532         45.932         -6.138         0.62           518         C         VAL         99         A         39.813         48.669         -4.532         0.52           519         O         VAL         99         A         40.679         48.871         -3.666         0.53           520         N         PRO         100         A         38.557         49.141         -4.403         0.53           521         CD         PRO         100         A         37.460         48.923         -5.352         0.54           522         CA         PRO         100         A         38.083         49.661         -3.125         0.49           523         CB         PRO         100         A         36.659         50.149         -3.411         0.50           524         CG         PRO         100         A         36.231         49.470         -4.666         0.47           525         C         PRO         100         A         37.913         47.407         -2.250         0.58           527         N         LYS         101         A				99	A	40.467	45.868	-4.573	0.44
518         C         VAL         99         A         39.813         48.669         -4.532         0.52           519         O         VAL         99         A         40.679         48.871         -3.666         0.53           520         N         PRO         100         A         38.557         49.141         -4.403         0.53           521         CD         PRO         100         A         37.460         48.923         -5.352         0.54           522         CA         PRO         100         A         38.083         49.661         -3.125         0.49           523         CB         PRO         100         A         36.659         50.149         -3.411         0.50           524         CG         PRO         100         A         36.231         49.470         -4.666         0.47           525         C         PRO         100         A         37.913         47.407         -2.250         0.58           527         N         LYS         101         A         37.739         49.029         -0.836         0.48           528         CA         LYS         101         A					A	38.532	45.932	-6.138	0.62
519       O       VAL       99       A       40.679       48.871       -3.666       0.53         520       N       PRO       100       A       38.557       49.141       -4.403       0.53         521       CD       PRO       100       A       37.460       48.923       -5.352       0.54         522       CA       PRO       100       A       38.083       49.661       -3.125       0.49         523       CB       PRO       100       A       36.659       50.149       -3.411       0.50         524       CG       PRO       100       A       36.231       49.470       -4.666       0.47         525       C       PRO       100       A       38.085       48.608       -2.041       0.41         526       O       PRO       100       A       37.913       47.407       -2.250       0.58         527       N       LYS       101       A       37.739       49.029       -0.836       0.48         528       CA       LYS       101       A       37.549       48.844       1.590       0.58         530       CG       LYS				99	A	39.813	48.669	-4.532	0.52
520         N         PRO         100         A         38.557         49.141         -4.403         0.53           521         CD         PRO         100         A         37.460         48.923         -5.352         0.54           522         CA         PRO         100         A         38.083         49.661         -3.125         0.49           523         CB         PRO         100         A         36.659         50.149         -3.411         0.50           524         CG         PRO         100         A         36.231         49.470         -4.666         0.47           525         C         PRO         100         A         38.085         48.608         -2.041         0.41           526         O         PRO         100         A         37.913         47.407         -2.250         0.58           527         N         LYS         101         A         37.739         49.029         -0.836         0.48           528         CA         LYS         101         A         37.549         48.844         1.590         0.58           530         CG         LYS         101         A				99	A	40.679	48.871	-3.666	0.53
522         CA         PRO 100         A         38.083         49.661         -3.125         0.49           523         CB         PRO 100         A         36.659         50.149         -3.411         0.50           524         CG         PRO 100         A         36.231         49.470         -4.666         0.47           525         C         PRO 100         A         38.085         48.608         -2.041         0.41           526         O         PRO 100         A         37.913         47.407         -2.250         0.58           527         N         LYS         101         A         37.739         49.029         -0.836         0.48           528         CA         LYS         101         A         37.365         48.136         0.241         0.40           529         CB         LYS         101         A         37.549         48.844         1.590         0.58           530         CG         LYS         101         A         38.756         48.257         2.367         0.69           531         CD         LYS         101         A         39.462         49.214         3.271         0.65 </td <td></td> <td></td> <td>PRO</td> <td>100</td> <td>A</td> <td>38.557</td> <td>49.141</td> <td>-4.403</td> <td>0.53</td>			PRO	100	A	38.557	49.141	-4.403	0.53
522         CA         PRO 100         A         38.083         49.661         -3.125         0.49           523         CB         PRO 100         A         36.659         50.149         -3.411         0.50           524         CG         PRO 100         A         36.231         49.470         -4.666         0.47           525         C         PRO 100         A         38.085         48.608         -2.041         0.41           526         O         PRO 100         A         37.913         47.407         -2.250         0.58           527         N         LYS 101         A         37.739         49.029         -0.836         0.48           528         CA         LYS 101         A         37.549         48.844         1.590         0.58           530         CG         LYS 101         A         38.756         48.257         2.367         0.69           531         CD         LYS 101         A         39.462         49.214         3.271         0.65           532         CE         LYS 101         A         39.223         51.248         4.589         0.76           533         NZ         LYS 101 </td <td>521</td> <td>CD</td> <td>PRO</td> <td>100</td> <td>A</td> <td>37.460</td> <td>48.923</td> <td>-5.352</td> <td>0.54</td>	521	CD	PRO	100	A	37.460	48.923	-5.352	0.54
523         CB         PRO 100         A         36.659         50.149         -3.411         0.50           524         CG         PRO 100         A         36.231         49.470         -4.666         0.47           525         C         PRO 100         A         38.085         48.608         -2.041         0.41           526         O         PRO 100         A         37.913         47.407         -2.250         0.58           527         N         LYS 101         A         37.739         49.029         -0.836         0.48           528         CA         LYS 101         A         37.365         48.136         0.241         0.40           529         CB         LYS 101         A         37.549         48.844         1.590         0.58           530         CG         LYS 101         A         38.756         48.257         2.367         0.69           531         CD         LYS 101         A         39.462         49.214         3.271         0.65           532         CE         LYS 101         A         38.839         50.616         3.297         0.61           533         NZ         LYS 101 <td></td> <td></td> <td>PRO</td> <td>100</td> <td>A</td> <td>38.083</td> <td>49.661</td> <td>-3.125</td> <td>0.49</td>			PRO	100	A	38.083	49.661	-3.125	0.49
524         CG         PRO         100         A         36.231         49.470         -4.666         0.47           525         C         PRO         100         A         38.085         48.608         -2.041         0.41           526         O         PRO         100         A         37.913         47.407         -2.250         0.58           527         N         LYS         101         A         37.739         49.029         -0.836         0.48           528         CA         LYS         101         A         37.365         48.136         0.241         0.40           529         CB         LYS         101         A         37.549         48.844         1.590         0.58           530         CG         LYS         101         A         38.756         48.257         2.367         0.69           531         CD         LYS         101         A         39.462         49.214         3.271         0.65           532         CE         LYS         101         A         38.839         50.616         3.297         0.61           533         NZ         LYS         101         A		CB	PRO	100	A	36.659	50.149	-3.411	0.50
525         C         PRO         100         A         38.085         48.608         -2.041         0.41           526         O         PRO         100         A         37.913         47.407         -2.250         0.58           527         N         LYS         101         A         37.739         49.029         -0.836         0.48           528         CA         LYS         101         A         37.365         48.136         0.241         0.40           529         CB         LYS         101         A         37.549         48.844         1.590         0.58           530         CG         LYS         101         A         38.756         48.257         2.367         0.69           531         CD         LYS         101         A         39.462         49.214         3.271         0.65           532         CE         LYS         101         A         38.839         50.616         3.297         0.61           533         NZ         LYS         101         A         39.223         51.248         4.589         0.76           534         C         LYS         101         A					A	36.231	49.470	-4.666	0.47
526         O         PRO         100         A         37.913         47.407         -2.250         0.58           527         N         LYS         101         A         37.739         49.029         -0.836         0.48           528         CA         LYS         101         A         37.365         48.136         0.241         0.40           529         CB         LYS         101         A         37.549         48.844         1.590         0.58           530         CG         LYS         101         A         38.756         48.257         2.367         0.69           531         CD         LYS         101         A         39.462         49.214         3.271         0.65           532         CE         LYS         101         A         38.839         50.616         3.297         0.61           533         NZ         LYS         101         A         39.223         51.248         4.589         0.76           534         C         LYS         101         A         35.850         47.825         0.048         0.56           535         O         LYS         101         A         <					A	38.085	48.608	-2.041	0.41
527         N         LYS         101         A         37.739         49.029         -0.836         0.48           528         CA         LYS         101         A         37.365         48.136         0.241         0.40           529         CB         LYS         101         A         37.549         48.844         1.590         0.58           530         CG         LYS         101         A         38.756         48.257         2.367         0.69           531         CD         LYS         101         A         39.462         49.214         3.271         0.65           532         CE         LYS         101         A         38.839         50.616         3.297         0.61           533         NZ         LYS         101         A         39.223         51.248         4.589         0.76           534         C         LYS         101         A         35.850         47.825         0.048         0.56           535         O         LYS         101         A         35.301         48.383         -0.922         0.54           536         N         PRO         102         A         <						37.913	47.407	-2.250	0.58
528         CA         LYS         101         A         37.365         48.136         0.241         0.40           529         CB         LYS         101         A         37.549         48.844         1.590         0.58           530         CG         LYS         101         A         38.756         48.257         2.367         0.69           531         CD         LYS         101         A         39.462         49.214         3.271         0.65           532         CE         LYS         101         A         38.839         50.616         3.297         0.61           533         NZ         LYS         101         A         39.223         51.248         4.589         0.76           534         C         LYS         101         A         35.850         47.825         0.048         0.56           535         O         LYS         101         A         35.301         48.383         -0.922         0.54           536         N         PRO         102         A         35.434         46.638         0.475         0.57           537         CD         PRO         102         A         <					A	37.739	49.029	-0.836	0.48
529         CB         LYS         101         A         37.549         48.844         1.590         0.58           530         CG         LYS         101         A         38.756         48.257         2.367         0.69           531         CD         LYS         101         A         39.462         49.214         3.271         0.65           532         CE         LYS         101         A         38.839         50.616         3.297         0.61           533         NZ         LYS         101         A         39.223         51.248         4.589         0.76           534         C         LYS         101         A         35.850         47.825         0.048         0.56           535         O         LYS         101         A         35.301         48.383         -0.922         0.54           536         N         PRO         102         A         35.434         46.638         0.475         0.57           537         CD         PRO         102         A         36.255         45.605         1.164         0.48           538         CA         PRO         102         A         <					A	37.365	48.136	0.241	0.40
530         CG         LYS         101         A         38.756         48.257         2.367         0.69           531         CD         LYS         101         A         39.462         49.214         3.271         0.65           532         CE         LYS         101         A         38.839         50.616         3.297         0.61           533         NZ         LYS         101         A         39.223         51.248         4.589         0.76           534         C         LYS         101         A         35.850         47.825         0.048         0.56           535         O         LYS         101         A         35.301         48.383         -0.922         0.54           536         N         PRO         102         A         35.434         46.638         0.475         0.57           537         CD         PRO         102         A         36.255         45.605         1.164         0.48           538         CA         PRO         102         A         34.017         46.290         0.428         0.59           539         CB         PRO         102         A         <			LYS	101	A	37.549	48.844	1.590	0.58
531         CD         LYS         101         A         39.462         49.214         3.271         0.65           532         CE         LYS         101         A         38.839         50.616         3.297         0.61           533         NZ         LYS         101         A         39.223         51.248         4.589         0.76           534         C         LYS         101         A         35.850         47.825         0.048         0.56           535         O         LYS         101         A         35.301         48.383         -0.922         0.54           536         N         PRO         102         A         35.434         46.638         0.475         0.57           537         CD         PRO         102         A         36.255         45.605         1.164         0.48           538         CA         PRO         102         A         34.017         46.290         0.428         0.59           539         CB         PRO         102         A         33.921         44.912         1.028         0.43		CG	LYS	101	A	38.756	48.257	2.367	0.69
532         CE         LYS 101         A         38.839         50.616         3.297         0.61           533         NZ         LYS 101         A         39.223         51.248         4.589         0.76           534         C         LYS 101         A         35.850         47.825         0.048         0.56           535         O         LYS 101         A         35.301         48.383         -0.922         0.54           536         N         PRO 102         A         35.434         46.638         0.475         0.57           537         CD         PRO 102         A         36.255         45.605         1.164         0.48           538         CA         PRO 102         A         34.017         46.290         0.428         0.59           539         CB         PRO 102         A         33.921         44.912         1.028         0.43		CD	LYS	101	A	39.462	49.214	3.271	0.65
533         NZ         LYS         101         A         39.223         51.248         4.589         0.76           534         C         LYS         101         A         35.850         47.825         0.048         0.56           535         O         LYS         101         A         35.301         48.383         -0.922         0.54           536         N         PRO         102         A         35.434         46.638         0.475         0.57           537         CD         PRO         102         A         36.255         45.605         1.164         0.48           538         CA         PRO         102         A         34.017         46.290         0.428         0.59           539         CB         PRO         102         A         33.921         44.912         1.028         0.43			LYS	101	A	38.839	50.616	3.297	0.61
534 C LYS 101 A 35.850 47.825 0.048 0.56 535 O LYS 101 A 35.301 48.383 -0.922 0.54 536 N PRO 102 A 35.434 46.638 0.475 0.57 537 CD PRO 102 A 36.255 45.605 1.164 0.48 538 CA PRO 102 A 34.017 46.290 0.428 0.59 539 CB PRO 102 A 33.921 44.912 1.028 0.43		NZ	LYS	101	A	39.223	51.248	4.589	0.76
535         O         LYS         101         A         35.301         48.383         -0.922         0.54           536         N         PRO         102         A         35.434         46.638         0.475         0.57           537         CD         PRO         102         A         36.255         45.605         1.164         0.48           538         CA         PRO         102         A         34.017         46.290         0.428         0.59           539         CB         PRO         102         A         33.921         44.912         1.028         0.43			LYS	101	A	35.850	47.825	0.048	0.56
536     N     PRO 102     A     35.434     46.638     0.475     0.57       537     CD     PRO 102     A     36.255     45.605     1.164     0.48       538     CA     PRO 102     A     34.017     46.290     0.428     0.59       539     CB     PRO 102     A     33.921     44.912     1.028     0.43					A	35.301	48.383	-0.922	0.54
537 CD PRO 102 A 36.255 45.605 1.164 0.48 538 CA PRO 102 A 34.017 46.290 0.428 0.59 539 CB PRO 102 A 33.921 44.912 1.028 0.43					A	35.434	46.638	0.475	0.57
538 CA PRO 102 A 34.017 46.290 0.428 0.59 539 CB PRO 102 A 33.921 44.912 1.028 0.43			PRO		A	36.255	45.605	1.164	0.48
539 CB PRO 102 A 33.921 44.912 1.028 0.43			PRO	102	A	34.017	46.290	0.428	0.59
27 222 14 772 4 727 2 7					A	33.921	44.912	1.028	0.43
	540	CG	PRO	102	A	35.209	44.663	1.736	0.54

72/98 **Fig. 16\13** 

A	tom	Residuc	Chain	x	Υ	Z	δ_
541	С	PRO 102	A	33.146	47.342	1.076	0.56
542	0	PRO 102	A	33.527	48.083	2.006	0.63
543	N	CYS 103	A	31.861	47.298	0.744	0.58
544	CA	CYS 103	A	30.857	48.177	1.400	0.48
545	CB	CYS 103	A	30.152	49.052	0.384	0.75
546	SG	CYS 103	A	28.673	48.328	-0.344	0.81
547	С	CYS 103	A	29.943	47.435	2.355	0.55
548	0	CYS 103	A	29.809	46.208	2.353	0.53
549	N	CYS 104	A	29.644	48.101	3.461	0.32
550	CA	CYS 104	A	28.717	47.709	4.468	0.48
551	С	CYS 104	A	27.265	47.729	3.963	0.60
552	0	CYS 104	A	26.729	48.760	3.570	0.60
553	CB	CYS 104	A	28.932	48.683	5.594	0.32
554	SG	CYS 104	A	28.146	48.223	7.124	0.54
555	N	ALA 105	λ	26.605	46.583	4.024	0.51
556	CA	ALA 105	A	25.412	46.300	3.222	0.51
557	CB	ALA 105	A	25.801	45.889	1.838	0.51
558	С	ALA 105	A	24.633	45.240	4.017	0.51
559	0	ALA 105	A	25.189	44.843	5.068	0.57
560	N	PRO 106	A	23.328	45.294	3.866	0.59
561	CD	PRO 106	A	22.564	46.026	2.822	0.51
562	CA	PRO 106	A	22.416	44.526	4.739	0.59
563	CB	PRO 106	A	21.014	45.021	4.321	0.54
564	CG	PRO 106	A	21.235	46.282	3.559	0.50
565	С	PRO 106	A	22.577	43.053	4.332	0.58
566	0	PRO 106	A	22.779	42.755	3.154	0.54
567	N	THR 107	A	22.616	42.182	5.306	0.60
568	CA	THR 107	A	22.791	40.733	4.913	0.65
569	CB	THR 107	A	23.831	40.113	5.928	0.45
570	OG1	THR 107	A	23.141	40.103	7.208	0.68
571	CG2	THR 107	A	25.153	40.852	6.091	0.56
572	С	THR 107	A	21.417	40.091	5.133	0.68
573	0	THR 107	A	21.037	39.069	4.543	0.68
574	N	GLN 108	A	20.597	40.806	5.902	0.67
575	CA	GLN 108	A	19.208	40.371	6.067	0.64
576	СВ	GLN 108	A	19.029	39.316	7.105	0.58
577	CG	GLN 108	A	19.016	39.718	8.547	0.70
578	CD	GLN 108	A	18.119	38.791	9.358	0.99
579	OE1	GLN 108	A	17.035	38.401	8.916	0.83
580	NE2	GLN 108	A	18.624	38.356	10.508	0.94
581	С	GLN 108	A	18.217	41.519	6.135	0.63
582	0	GLN 108	A	18.323	42.360	7.047	0.64
583	N	LEU 109	A	17.213	41.401	5.288	0.60
584	CA	LEU 109	A	15.994	42.229	5.307	0.65
585	СВ	LEU 109	A	15.882	42.694	3.831	0.55

73/98 *Fig. 16\14* 

A	lom	Residue	Chain	x	Y	z	δ
586	CG	LEU 109	A	17.101	43.501	3.407	0.59
587	CD1	LEU 109	A	17.359	43.341	1.936	0.68
588	CD2	LEU 109	A	16.896	44.975	3.746	0.56
589	C	LEU 109	A	14.740	41.492	5.774	0.70
590	0	LEU 109	A	14.393	40.460	5.142	0.75
591	N	ASN 110	A	13.876	42.190	6.507	0.67
592	CA	ASN 110	A	12.512	41.767	6.849	0.65
593	CB	ASN 110	A	12.192	41.773	8.350	0.59
594	CG	ASN 110	A	13.043	40.783	9.115	0.79
595	OD1	ASN 110	· <b>A</b>	13.197	40.850	10.340	0.90
596	ND2	ASN 110	A	13.591	39.872	8.302	0.79
597	С	ASN 110	A	11.441	42.589	6.116	0.65
598	0	ASN 110	A	11.655	43.703	5.641	0.61
599	N	ALA 111	A	10.255	41.992	6.127	0.68
600	CA	ALA 111	A	9.062	42.537	5.489	0.64
601	СВ	ALA 111	A	8.158	41.466	4.906	0.59
602	С	ALA 111	A	8.315	43.435	6.467	0.57
603	0	ALA 111	A	8.500	43.354	7.697	0.54
604	N	ILE 112	A	7.548	44.344	5.879	0.54
605	CA	ILE 112	A	6.518	45.030	6.743	0.47
606	CB	ILE 112	A	6.821	46.531	6.985	0.63
607	CG2	ILE 112	A	7.990	46.879	7.923	0.68
608	CG1	ILE 112	A	6.825	47.379	5.712	0.56
609	CD	ILE 112	A	6.369	48.858	5.888	0.64
610	С	ILE 112	A	5.183	44.777	5.997	0.45
611	0	ILE 112	A	5.166	44.766	4.747	0.54
612	N	SER 113	A	4.150	44.802	6.770	0.44
613	CA	SER 113	A	2.734	44.868	6.330	0.61
614	CB	SER 113	A	1.946	43.791	7.102	0.63
615	OG	SER 113	A	2.163	42.503	6.515	0.62
616	С	SER 113	Α	2.172	46.244	6.694	0.53
617	0	SER 113	A	2.299	46.730	7.828	0.52
618	N	VAL 114	A	1.400	46.804	5.778	0.55
619	CA	VAL 114	A	0.651	48.025	6.121	0.58
620	СВ	VAL 114	A	1.371	49.244	5.514	0.64
621	CG1	VAL 114	A	2.885	49.190	5.667	0.49
622	CG2	VAL 114	A	0.949	49.424	4.094	0.55
623	C	VAL 114	A	-0.813	47.879	5.686	0.60
624	0	VAL 114	A	-1.189	47.327	4.637	0.51
625	N	LEU 115	A	-1.652	48.484	6.510	0.49
626	CA	LEU 115	A	-3.116	48.419	6.182	0.58
627	CB	LET 115	A	-3.789	48.328	7.603	0.55
628	CG	LEU 115	A	-5.284	47.999	7.568	0.57
629	CD1	LEU 115	A	-5.523	46.603	7.036	0.59
630	CD2	LEU 115	A	-5.843	48.190	8.975	0.52

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Fig. 16\15

A	om	Residuc	Chain	X	Y	Z	δ
			_		40 605	F 400	A 45
631	Ç	LEU 115	A	-3.421	49.685	5.403	0.47
632	0	LEU 115	A	-3.244	50.761	5.955	0.57
633	N	TYR 116	A	-3.971	49.616	4.196	0.57
634	CA	TYR 116	A	-4.302	50.847	3.479	0.69
635	CB	TYR 116	A	-3.192	51.211	2.480	0.70
636	CG	TYR 116	A	-3.119	50.316	1.269	0.46
637	CD1	TYR 116	A	-2.487	49.097	1.272	0.57
638	CE1	TYR 116	A	-2.542	48.256	0.154	0.66
639	CD2	TYR 116	A	-3.584	50.775	0.050	0.57
640	CE2	TYR 116	A	-3.689	49.971	-1.067	0.52
641	CZ	TYR 116	A	-3.151	48.697	-1.016	0.76
642	OH	TYR 116	A	-3.119	47.979	-2.184	0.78
643	С	TYR 116	A	-5.692	50.766	2.849	0.69
644	0	TYR 116	A	-6.250	49.665	2.752	0.69
645	N	PHE 117	A	-6.209	51.919	2.485	0.56
646	CA	PHE 117	A	-7.443	52.158	1.793	0.60
647	CB	PHE 117	A	-8.462	53.064	2.413	0.64
648	CG	PHE 117	A	-8.074	54.254	3.206	0.B2
649	CD1	PHE 117	A	-8.221	54.256	4.599	0.91
650	CD2	PHE 117	A	-7.813	55.472	2.561	1.11
651	CE1	PHE 117	A	-7.964	55.401	5.343	1.12
652	CE2	PHE 117	A	-7.547	56.638	3.292	1.03
653	CZ	PHE 117	A	-7.638	56.601	4.697	1.02
654	C	PHE 117	A	-7.240	52.362	0.315	0.62
<b>65</b> 5	0	PHE 117	A	-6.666	53.381	-0.059	0.72
656	N	ASP 118	A	-7.607	51.365	-0.474	0.67
657	CA	ASP 118	A	-7.388	51.415	-1.922	0.67
658	CB	ASP 118	A	-7.342	50.074	-2.610	0.73
659	CG	ASP 118	A	-8.694	49.558	-3.079	0.82
660	OD1	ASP 118	A	-8.748	48.554	-3.798	0.72
661	OD2	ASP 118	A	-9.705	50.249	-2.831	0.60
662	¢	ASP 118	A	-8.337	52.433	-2.526	0.68
663	0	ASP 118	A	-9.108	53.096	-1.826	0.75
664	N	ASP 119	A	-8.344	52.429	-3.851	0.76
665	CA	ASP 119	A	-9.035	53.486	-4.614	0.83
666	CB	ASP 119	A	-8.356	53.711	-5.961	0.94
667	CG	ASP 119	A	-7.989	52.393	-6.634	1.08
668	OD1	ASP 119	A	-8.883	51.605	-6.991	1.04
669	OD2	ASP 119	A	-6.773	52.103	-6.627	1.06
670	С	ASP 119	A	-10.531	53.204	-4.673	0.87
671	0	ASP 119	A	-11.339	54.138	-4.854	0.90
672	N	SER 120	A	-10.886	51.946	-4.448	0.85
673	CA	SER 120	A	-12.314	51.574	-4.389	0.80
674	CB	SER 120	A	-12.618	50.258	-5.030	0.83
675	OG	SER 120	A	-11.534	49.729	-5.781	1.02

75/98 **Fig. 16\16** 

At	om	Residue	Chain	x	Y	Z	8
	_	ann 120	A	-12.796	51.719	-2.957	0.80
676	C	SER 120	A	-14.013	51.770	-2.686	0.83
677	0	SER 120	A	-11.823	51.902	-2.064	0.73
678	N	SER 121 SER 121	λ	-12.197	52.163	-0.653	0.69
679	CA	SER 121 SER 121	A	-13.578	52.816	-0.607	0.66
680	CB	SER 121	A	-13.479	54.163	-1.016	0.85
681	oG	SER 121	A	-12.186	50.879	0.163	0.63
682	C	SER 121	A	-12.564	50.874	1.332	0.69
683	0	ASN 122	A	-11.745	49.825	-0.477	0.65
684	N	ASN 122	A	-11.388	48.578	0.218	0.63
685	CA CB	ASN 122	A	-11.103	47.558	-0.897	0.69
686	CG	ASN 122	Α	-12.211	47.680	-1.955	0.73
687	OD1	ASN 122	A	-13.370	47.437	-1.5 <del>9</del> 7	0.73
688 689	ND2	ASN 122	A	-11.867	47.636	-3.240	0.75
690	C	ASN 122	A	-10.237	48.837	1.176	0.67
691	0	ASN 122	A	-9.323	49.604	0.852	0.63
692	N	VAL 123	A	-10.305	48.227	2.346	0.66
693	CA	VAL 123	A	-9.252	48.270	3.360	0.61
694	СВ	VAL 123	A	-9.927	48.279	4.744	0.58
695	CG1	VAL 123	A	-8.929	47.747	5.758	0.59
696	CG2	VAL 123	A	-10.405	49.677	5.089	0.56
697	C	VAL 123	A	-8.351	47.044	3.220	0.63
698	0	VAL 123	A	-8.901	45.942	3.098	0.65
699	N	ILE 124	A	-7.133	47.256	2.758	0.70
700	CA	ILE 124	A	-6.224	46.189	2.302	0.65
701		ILE 124	A	-5.819	46.559	0.813	0.67
702		ILE 124	A	-4.839	45.531	0.198	0.67
703		ILE 124	A	-7.136	46.616	0.011	0.66
704		ILE 124	A	-7.044	46.191	-1.468	0.84
705		ILE 124	A	-4.998	46.057	3.208	0.67
706	0	ILE 124	Α	-4.326	47.034	3.582	0.63
707	N	LEU 125	A	-4.598	44.815	3.448	0.69
708	CA	LET 125	A ·	-3.323	44.541	4.140 5.187	0.67
709	CB	LEU 125	A	-3.602	43.477	6.067	0.64
710	CG	LEU 125	A	-2.441	43.036 44.105	7.130	0.49
711	CD1	LEU 125	A	-2.163	41.760	6.750	0.48
712		LEU 125	A	-2.977 -2.249	44.132	3.135	0.67
713		LEU 125	A	-2.249 -2.431	43.103	2.470	0.66
714		LEU 125	A	-1.263	44.993	2.961	0.66
715		LYS 126	A	-0.174	44.777	2.001	0.59
716		LYS 126	A	-0.174	45.640	0.788	0.61
717		LYS 126	Α	1.066		-0.128	0.78
718		LYS 126	A	0.687	45.478	-1.601	0.91
719		LYS 126	A	1.833		-2.543	0.82
720	CE	LYS 126	A	1.633	-5.25		

<sup>76/98</sup> *Fig. 16\17* 

A	tom	Residue	Chain	x	Y	Z	δ
721	NZ	LYS 126	A	1.346	44.434	-3.746	0.86
722	C	LYS 126	À	1.173	44.619	2.669	0.50
723	0	LYS 126	A	1.398	44.978	3.845	0.57
724	N	LYS 127	A	1.836	43.565	2.204	0.61
725	CA	LYS 127	A	3.124	43.060	2.664	0.60
726	СВ	LYS 127	A	3.198	41.534	2.620	0.58
727	CG	LYS 127	A	4.330	40.965	3.471	0.57
728	CD	LYS 127	A	4.828	39.671	2.833	0.62
729	CE	LYS 127	A	5.639	38.881	3.845	0.76
730	NZ	LYS 127	A	5.283	39.360	5.208	0.83
731	C	LYS 127	A	4.245	43.589	1.755	0.60
732	0	LYS 127	A	4.058	43.699	0.521	0.49
733	N	TYR 128	A	5.136	44.302	2.424	0.56
734	CA	TYR 128	A	6.222	44.975	1.669	0.50
735	CB	TYR 128	A	6.302	46.457	2.089	0.77
736	CG	TYR 128	A	5.310	47.284	1.298	0.64
737	CD1	TYR 128	A	4.269	47.954	1.935	0.68
738	CE1	TYR 128	A	3.320	48.646	1.174	0.68
739	CD2	TYR 128	A	5.280	47.163	-0.085	0.72
740	CE2	TYR 128	A	4.318	47.820	-0.857	0.77
741	CZ	TYR 128	A	3.268	48.459	-0.200	0.89
742	OH	TYR 128	A	2.279	49.021	-0.961	0.90
743	С	TYR 128	A	7.527	44.284	2.110	0.46
744	0	TYR 128	A	7.781	44.139	3.308	0.59
745	N	ARG 129	A	7.978	43.511	1.150	0.56
746	CA	ARG 129	A	9.194	42.698	1.396	0.61
747	CB	ARG 129	A	9.175	41.646	0.273	0.74
748	CG	ARG 129	A	8.115	40.550	0.447	0.74
749	CD	ARG 129	A	8.486	39.314	-0.337	0.61
750	NE	ARG 129	A	9.295	38.414	0.488	1.06
751	CZ	ARG 129	A	10.599	38.183	0.310	1.07
752	NH1	ARG 129	A	11.211	38.363	-0.861	0.91
753	NH2	ARG 129	A	11.370	37.849	1.349	0.87
754	С	ARG 129	A	10.405	43.636	1.302	0.54
<b>75</b> 5	0	ARG 129	A	10.406	44.605	0.522	0.55
756	N	ASN 130	A	11.403	43.372	2.095	0.60
757	CA	ASN 130	A	12.792	43.856	2.013	0.59
758	CB	ASN 130	A	13.359	43.611	0.610	0.52
759	CG	ASN 130	A	13.760	42.157	0.392	0.63
760	001	ASN 130	A	14.053	41.398	1.318	0.58
761	ND2	ASN 130	A	13.517	41.609	-0.782	0.55
762	C	ASN 130	A	12.807	45.343	2.380	0.55
763	0	ASN 130	A	13.426	46.094	1.648	0.48
764	N	MET 131	A	12.267	45.671	3.520	0.54
765	CA	MET 131	A	11.990	47.026	3.974	0.51

77/98 *Fig. 16\18* 

A	tom	Residue	Chain	<u> </u>	Y		δ
766	CB	MET 131	A	10.470	47.189	4.327	0.54
767	CG	MET 131	A	9.590	47.290	3.137	0.44
768	SD	MET 131	A	9.927	48.720	2.052	0.61
769	CE	MET 131	A	8.873	49.928	2.915	0.60
770	С	MET 131	A	12.790	47.266	5.256	0.52
771	0	MET 131	A	13.048	48.429	5.570	0.59
772		VAL 132	A	12.997	46.230	6.055	0.59
773	CA	VAL 132	A	13.670	46.399	7.359	0.60
774	CB	VAL 132	A	12.807	46.001	8.517	0.52
775	CG1	VAL 132	A	13.412	46.036	9.898	0.48
776	CG2	VAL 132	A	11.331	46.318	8.485	0.66
777	C	VAL 132	A	15.035	45.704	7.308	0.69
778	0	VAL 132	A	15.148	44.495	7.012	0.63
779	N	VAL 133	A	16.039	46.488	7.668	0.65
780	CA	VAL 133	A	17.424	45.971	7.709	0.62
781	CB	VAL 133	A	18.486	47.081	7.591	0.56
782	CG1	VAL 133	A	19.842	46.568	8.105	0.53
783	CG2	VAL 133	A	18.577	47.505	6.128	0.39
784	C	VAL 133	A	17.542	45.241	9.052	0.62
785	0	VAL 133	A	17.392	45.958	10.044	0.54
786	N	ARG 134	A	17.821	43.930	8.941	0.53
787	CA	ARG 134	A	17.924	43.188	10.208	0.56
788	CB	ARG 134	A	17.114	41.877	10.158	0.68
789	CG	ARG 134	A	15.779	41.942	10.908	0.80
790	CD	ARG 134	A	15.932	41.805	12.379	0.79
791	NE	ARG 134	A	15.072	40.814	12.976	1.02
792	CZ	ARG 134	A	15.265	39.514	13.159	1.10
793	NH 1	ARG 134	A	16.185	38.783	12.533	1.08
794	NH2	ARG 134	A	14.403	38.858	13.952	1.09
795	С	ARG 134	A	19.370	42.957	10.622	0.57
796	0	ARG 134	A	19.660	42.742	11.813	0.61
797	N	ALA 135	A	20.282	42.931	9.659	0.53
798	CA	ALA 135	A	21.722	42.782	9.986	0.49
799	CB	ALA 135	λ	22.059	41.272	9.934	0.64
800	С	ALA 135	A	22.466	43.400	8.774	0.42
801	0	ALA 135	A	21.937	43.282	7.673	0.49
802	N	CYS 136	A	23.726	43.635	9.022	0.59
803	CA	CYS 136	A	24.682	44.274	8.089	0.67
804	C	CYS 136	A	25.954	43.443	7.969	0.61
805	0	CYS 136	A	26.383	42.850	8.985	0.59
806	CB	CYS 136	A	25.073	45.641	8.701	0.61
807	SG	CYS 136	A	23.632	46.772	8.673	0.61
808	N	GLY 137	A	26.579	43.476	6.808	0.59
809	CA	GLY 137	A	27.915	42.790	6.772	0.59
810	С	GLY 137	λ	28.804	43.523	5.787	0.52

78/98 **Fig. 16\19** 

At	om	Res	iduc	Chain	X	Y	z	δ
811	0	GLY	137	A	28.249	44.112	4.845	0.49
812	И	_	138	A	30.019	43.036	5.634	0.57
813	CA		138	A	30.912	43.537	4.545	0.48
814	c	CYS	138	A	30.845	42.606	3.369	0.56
815	0		138	A	31.335	41.457	3.446	0.55
816	СВ		138	A	32.348	43.567	5.123	0.38
817	SG		138	A	32.401	44.877	6.428	0.57
818	N		139	A	30.522	43.184	2.210	0.63
819	CA		139	A	30.557	42.395	0.970	0.54
820	CB		139	A	29,665	41.188	0.842	0.88
821	CG	HIS		A	28.437	41.289	1.707	0.94
822	CD2	HIS		A	27.571	42.337	1.811	0.79
823	ND1	HIS		A	28.113	40.406	2.706	0.91
B24	CE1	HIS	139	A	27.001	40.825	3.288	0.97
825	NE2	HIS	139	A	26.665	41.993	2.774	0.96
826	С	HIS	139	A	30.936	43.135	-0.264	0.58
827	OT1	HIS	139	A	31.164	42.426	~1.268	0.75
828	OT2	HIS	139	A	31.123	44.371	-0.256	0.79
829	CB	GLN	36	В	29.653	57.175	-11.979	0.80
830	CG	GLN	36	В	30.943	57.649	-11.327	0.98
831	CD	GLN	36	В	30.751	58.560	-10.142	1.07
832	OE1	GLN	36	В	29.879	59.426	-10.061	0.98
833	NE2	GLN	36	В	31.585	58.331	-9.123	1.05
834	С	GLN	36	В	28.572	54.939	-11.583	0.62
835	0	GLN	36	В	28.641	54.928	-10.349	0.67
836	N	GLN	36	В	29.482	55.414	-13.781	0.82
837	CA	GLN	36	В	29.627	55.707	-12.360	0.72
838	N	ALA	37	В	27.621	54.379	-12.314	0.65
839	CA	ALA	37	В	26.512	53.646	-11.644	0.64
840	CB	ALA	37	B	25.331	53.455	-12.576	0.58
841	С	ALA	37	В	27.011	52.266	-11.201	0.62
842	0	ALA	37	В	27.851	51.674	-11.868	0.67
843	N	CYS	38	В	26.260	51.702	-10.285	0.64
844	CA	CYS	38	В	26.531	50.390	-9.714	0.53
845	C	CYS	38	В	26.592	49.339	-10.797	0.63
846	0	CYS	38	В	25.623	49.141	-11.527	0.66
847	CB	CYS	38	В	25.607	50.071	-8.582	0.58
848	SG	CYS	38	В	25.827	48.462	-7.828	0.63
849	N	LYS	39	В	27.800		-11.010	0.63
850	CA	LYS	39	В	28.056		-11.860	0.50
851	CB	LYS	39	В	28.442		-13.296	0.56
852	CG	LYS	39	В	29.121		-13.549	0.69
853	CD	LYS	39	В	29.867		-14.870	0.62
854	CE	LYS	39	В	29.840		-15.592	0.92
855	NZ	LYS	39	В	30.922	47.935	-16.607	0.92

79/98 **Fig. 16\20** 

857 O LYS 39 B 29.788 47.073 -10.242   858 N LYS 40 B 29.405 45.717 -11.988   859 CA LYS 40 B 30.309 44.615 -11.659   860 CB LYS 40 B 30.617 42.119 -11.621   861 CG LYS 40 B 30.617 42.119 -11.621   862 CD LYS 40 B 30.264 40.554 -13.565   863 CE LYS 40 B 30.264 40.554 -13.565   864 NZ LYS 40 B 30.264 40.554 -13.565   865 C LYS 40 B 31.570 44.807 -12.504   866 C LYS 40 B 31.570 44.807 -12.504   866 C LYS 40 B 31.450 45.250 -13.659   867 N HIS 41 B 32.700 44.888 -11.826   868 CA HIS 41 B 33.938 45.171 -12.590   869 CB HIS 41 B 33.634 45.771 -12.590   870 CG HIS 41 B 33.564 48.557 -12.835   871 CD2 HIS 41 B 32.931 47.989 -10.838   872 ND1 HIS 41 B 32.837 49.537 -12.338   873 CE1 HIS 41 B 32.837 49.537 -12.338   874 NE2 HIS 41 B 32.837 49.537 -12.338   875 C HIS 41 B 32.837 49.537 -12.338   876 O HIS 41 B 32.837 49.537 -12.338   877 N GLU 42 B 35.819 43.974 -13.373   878 CA GLU 42 B 36.805 42.897 -13.460   879 CB GLU 42 B 36.805 42.897 -13.460   879 CB GLU 42 B 36.806 41.769 -17.171   882 OE1 GLU 42 B 36.806 41.769 -17.171   883 OE2 GLU 42 B 36.806 41.769 -17.171   886 CB LEU 43 B 39.100 42.016 -9.561   888 CB LEU 43 B 39.970 42.470 -8.455   889 CG LEU 43 B 39.970 42.470 -8.455   899 CD1 LEU 43 B 39.970 42.470 -8.455   899 CD1 TYR 44 B 44.404 36.795 -12.620   899 CD1 TYR 44 B 44.404 36.795 -12.742   899 CD1 TYR 44 B 44.404 36.795 -1	LYS 39 B 29.788 47.073 -10.242 0.4 LYS 40 B 29.405 45.717 -11.988 0.5 LYS 40 B 29.713 43.258 -12.132 0.6 LYS 40 B 30.309 44.615 -11.659 0.6 LYS 40 B 30.617 42.119 -11.621 0.6 LYS 40 B 30.087 40.791 -12.085 0.4 LYS 40 B 30.264 40.554 -13.565 0.6 LYS 40 B 30.264 40.554 -13.565 0.6 LYS 40 B 30.264 40.554 -13.565 0.6 LYS 40 B 31.570 44.807 -12.504 0.6 LYS 40 B 31.570 44.807 -12.504 0.6 LYS 40 B 31.570 44.807 -12.504 0.6 LYS 40 B 31.570 44.807 -12.500 0.5 LYS 40 B 31.570 44.807 -12.500 0.6 LYS 40 B 31.570 44.807 -12.500 0.5 LYS 40 B 31.570 44.807 -12.500 0.6 LYS 40 B 31.570 44.807 -12.500 0.6 LYS 40 B 31.570 44.807 -12.500 0.6 LYS 40 B 31.570 44.808 -11.826 0.6 LYS 40 B 31.570 44.807 -12.500 0.5 LYS 40 B 31.570 44.808 -11.826 0.6 LYS 40 B 31.570 44.807 -12.500 0.5 LYS 40 B 31.570 -12.600 0.5 LYS 40 B 31.570 -12.600 0.5 LYY 41 B 31.500 43.777 -11.500 0.5 LY 41 B 31.500 43.777 -11.500 0.5 LY 42 B 38.304 41.805 -11.500 0	Atom		Resid	lue	Chain	x	Y	Z	δ
## 857 O LYS 39 B 29.788 47.073 -10.242   ## 858 N LYS 40 B 29.405 45.717 -11.988   ## 859 CA LYS 40 B 30.309 44.615 -11.659   ## 860 CB LYS 40 B 30.087 40.791 -12.085   ## 861 CG LYS 40 B 30.087 40.791 -12.085   ## 862 CD LYS 40 B 30.087 40.791 -12.085   ## 863 CE LYS 40 B 30.264 40.554 -13.565   ## 864 NZ LYS 40 B 30.264 40.554 -13.659   ## 865 C LYS 40 B 31.570 44.807 -12.504   ## 866 C LYS 40 B 31.570 44.807 -12.504   ## 866 C LYS 40 B 31.570 44.808 -11.826   ## 867 N HIS 41 B 32.700 44.888 -11.826   ## 868 CA HIS 41 B 33.938 45.171 -12.590   ## 869 CB HIS 41 B 33.639 47.549 -11.899   ## 870 CG HIS 41 B 33.639 47.549 -11.899   ## 871 CD2 HIS 41 B 32.931 47.989 -10.838   ## 872 ND1 HIS 41 B 32.931 47.989 -10.838   ## 873 CE1 HIS 41 B 32.837 49.537 -12.338   ## 874 NE2 HIS 41 B 32.837 49.537 -12.338   ## 875 C HIS 41 B 34.837 43.952 -12.505   ## 876 C HIS 41 B 34.837 43.952 -12.505   ## 877 N GLU 42 B 36.805 42.897 -13.460   ## 879 CB GLU 42 B 36.805 42.897 -13.460   ## 879 CB GLU 42 B 36.806 41.769 -17.171   ## 880 CG GLU 42 B 36.806 41.769 -17.171   ## 881 CD GLU 42 B 36.806 41.769 -17.171   ## 882 OE1 GLU 42 B 36.806 41.769 -17.171   ## 888 CB LEU 43 B 39.970 42.470 -8.455   ## 889 CB LEU 43 B 39.970 42.470 -8.455   ## 899 CB TYR 44 B 41.800 43.679 -11.590   ## 899 CB TYR 44 B 44.762 35.449 -11.570   ## 899 CB TYR 44 B 44.762 35.449 -11.574   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 44 B 44.762 35.449 -12.742   ## 899 CE1 TYR 4	LYS   39   B   29.788   47.073   -10.242   0.4									
858 N LYS 40 B 30.309 44.615 -11.659 860 CB LYS 40 B 30.309 44.615 -11.659 861 CG LYS 40 B 30.617 42.119 -11.621 862 CD LYS 40 B 30.617 42.119 -11.621 862 CD LYS 40 B 30.264 40.554 -13.565 863 CE LYS 40 B 30.264 40.554 -13.565 864 NZ LYS 40 B 30.264 40.554 -13.873 865 C LYS 40 B 31.570 44.807 -12.504 866 C LYS 40 B 31.450 45.250 -13.659 867 N HIS 41 B 32.700 44.808 -11.826 868 CA HIS 41 B 33.938 45.171 -12.590 869 CB HIS 41 B 33.639 47.549 -11.899 871 CD2 HIS 41 B 33.639 47.549 -11.899 871 CD2 HIS 41 B 32.837 47.989 -10.838 873 CE1 HIS 41 B 32.837 49.537 -12.338 874 NE2 HIS 41 B 32.837 49.537 -12.338 874 NE2 HIS 41 B 32.837 49.537 -12.338 875 C HIS 41 B 32.837 49.537 -12.338 876 O HIS 41 B 32.837 49.537 -12.338 877 N GLU 42 B 35.819 43.974 -13.373 878 CA GLU 42 B 35.819 43.974 -13.373 878 CA GLU 42 B 36.805 42.897 -13.460 879 CB GLU 42 B 36.805 42.897 -13.460 879 CB GLU 42 B 36.805 42.897 -13.460 889 CE GLU 42 B 36.806 41.769 -17.171 888 CB LEU 43 B 39.504 41.895 -11.858 890 CD1 LEU 43 B 39.120 42.016 -9.561 889 CB LEU 43 B 39.120 42.016 -9.561 889 CB LEU 43 B 39.970 42.470 -8.455 899 CD1 LEU 43 B 39.970 42.470 -8.455 899 CD1 TYR 44 B 41.804 36.795 -12.284 899 CE1 TYR 44 B 44.404 36.795 -12.284 899 CE1 TYR 44 B 44.404 36.795 -12.284 899 CE1 TYR 44 B 44.404 36.795 -12.242 899 CE1 TYR 44 B 44.404 36.795 -12.284 899 CE1 TYR 44 B 44	EXIS 40 B 29.405 45.717 -11.988 0.5  CA LYS 40 B 30.309 44.615 -11.659 0.6  CB LYS 40 B 30.617 42.119 -11.621 0.6  CD LYS 40 B 30.617 42.119 -11.621 0.6  CD LYS 40 B 30.617 42.119 -11.621 0.6  CD LYS 40 B 30.624 40.554 -13.565 0.6  CE LYS 40 B 30.264 40.554 -13.565 0.6  CE LYS 40 B 30.264 40.554 -13.565 0.6  CE LYS 40 B 31.570 44.807 -12.504 0.6  CE LYS 40 B 31.570 44.807 -12.505 0.6  CE HIS 41 B 32.931 47.989 -10.838 0.6  CE HIS 41 B 32.837 49.537 -12.335 0.6  CE HIS 41 B 32.837 49.537 -12.336 0.6  CE HIS 41 B 32.837 49.537 -12.336 0.6  CE HIS 41 B 32.837 49.537 -12.336 0.6  CE HIS 41 B 34.737 43.143 -11.564 0.6  CE HIS 41 B 34.737 43.143 -11.564 0.6  CE HIS 41 B 34.737 43.143 -11.564 0.6  CE GLU 42 B 36.806 42.897 -13.460 0.6  CE GLU 42 B 36.806 41.769 -17.171 0.8  CE GLU 42 B 38.361 44.183 -12.261 0.6  CE LEU 43 B 39.500 34.57 -0.7043 0.6  CE LEU 43 B 39.970 42.470 -8.455 0.6  CE LEU 43 B 39.970 42.470 -8.455 0.6  CE LEU 43 B 39.970 42.470 -8.455 0.6  CE LEU 43 B 39.947 40.315 -11.210 0.6  CE LEU 43 B 39.947 40.31	<b>B</b> 56	С	LYS	39	В				
859 CA LYS 40 B 29.713 43.258 -12.132 861 CG LYS 40 B 29.713 43.258 -12.132 861 CG LYS 40 B 30.617 42.119 -11.621 862 CD LYS 40 B 30.264 40.554 -13.565 863 CE LYS 40 B 30.264 40.554 -13.565 864 NZ LYS 40 B 31.570 44.807 -12.504 865 C LYS 40 B 31.570 44.807 -12.504 866 O LYS 40 B 31.570 44.808 -13.659 867 N HIS 41 B 32.700 44.808 -11.826 868 CA HIS 41 B 33.938 45.171 -12.590 869 CB HIS 41 B 33.639 47.549 -11.899 871 CD2 HIS 41 B 33.639 47.549 -11.899 871 CD2 HIS 41 B 32.931 47.989 -10.838 872 ND1 HIS 41 B 32.837 49.537 -12.338 874 NE2 HIS 41 B 32.837 49.537 -12.338 874 NE2 HIS 41 B 32.274 49.130 -11.217 875 C HIS 41 B 32.274 49.130 -11.217 875 C HIS 41 B 34.837 43.952 -12.505 876 O HIS 41 B 34.837 43.952 -12.505 876 O HIS 41 B 34.837 43.952 -12.505 876 O HIS 41 B 34.837 43.952 -12.505 876 CA GLU 42 B 35.819 43.974 -13.373 878 CA GLU 42 B 35.819 43.974 -13.373 878 CA GLU 42 B 36.805 42.897 -13.460 879 CB GLU 42 B 36.806 41.769 -17.171 882 OE1 GLU 42 B 38.361 44.883 -12.261 888 CB LEU 43 B 39.970 42.470 -8.455 888 CB LEU 43 B 39.970 42.470 -8.455 889 CD LEU 43 B 39.970 42.470 -8.455 889 CD LEU 43 B 39.970 42.470 -8.455 899 CD LEU 43 B 39.947 40.315 -11.210 899 CD LEU 43 B 39.970 42.470 -8.455 899 CD LEU 43 B 39.947 40.315 -11.210 899 CD LEU 43 B 39.947 40.315 -11.910 899 CD LEU 43 B 39.947 40.315 -11.91	TYS 40 B 30.309 44.615 -11.659 0.6  TYS 40 B 30.309 44.615 -11.659 0.6  TYS 40 B 30.617 42.119 -11.621 0.6  TYS 40 B 30.087 40.791 -12.085 0.6  TYS 40 B 30.087 40.791 -12.085 0.6  TYS 40 B 30.264 40.554 -13.565 0.6  TYS 40 B 30.264 40.554 -13.565 0.6  TYS 40 B 31.570 44.807 -12.504 0.6  TYS 40 B 31.570 44.807 -12.504 0.6  TYS 40 B 31.570 44.807 -12.504 0.6  TYS 40 B 31.570 44.808 -11.826 0.6  TYS 40 B 31.570 45.250 -13.659 0.6  TYS 40 B 31.570 45.250 -13.659 0.6  TYS 40 B 31.570 45.250 -13.659 0.6  TYS 40 B 31.570 45.250 -12.505 0.6  TYS 40 B 31.570 45.250 -12.505 0.6  TYS 40 B 31.570 44.808 -11.826 0.6  TYS 41 B 32.931 47.989 -10.838 0.6  TYS 41 B 32.931 47.989 -10.838 0.6  TYS 41 B 32.837 49.537 -12.338 0.6  TYS 41 B 32.837 49.537 -12.338 0.6  TYS 41 B 32.274 49.130 -11.217 0.6  TYS 41 B 34.837 43.952 -12.505 0.5  TYS 40 B 36.806 41.769 -17.171 0.6  TYS 40 B 36.806 41.769 -17.171 0.6  TYS 40 B 39.507 42.470 -8.455 0.6  TYS 40 B 39.538 41.815 -11.012 0.5  TYS 40 B 39.947 40.315 -11.210 0.6  TYS 44 B 41.300 43.057 -8.715 0.6  TYS 44 B 41.300 43.057 -8.715 0.6  TYS 44 B 41.300 43.057 -8.715 0.6  TYS 44 B 41.300 43.057 -11.580 0.6  TYS 44 B 41.300 43.057 -11.580 0.6  TYS 44 B 41.404 36.795 -12.620 0.6  TYS 44 B 44.662 35.449 -12.742 0.6  TYS 44 B 44.604 36.795 -12.620 0.6	857	0	LYS	39	В				
860 CB LYS 40 B 30.617 42.119 -11.621 862 CD LYS 40 B 30.617 42.119 -11.621 862 CD LYS 40 B 30.087 40.791 -12.085 863 CE LYS 40 B 30.264 40.554 -13.565 864 NZ LYS 40 B 31.570 44.807 -12.504 866 C LYS 40 B 31.570 44.807 -12.504 866 C LYS 40 B 31.570 44.807 -12.504 867 N HIS 41 B 32.700 44.808 -11.826 868 CA HIS 41 B 33.938 45.171 -12.590 869 CB HIS 41 B 33.639 47.549 -11.899 871 CD2 HIS 41 B 32.931 47.989 -10.838 872 ND1 HIS 41 B 32.931 47.989 -10.838 873 CE1 HIS 41 B 32.837 49.537 -12.338 874 NE2 HIS 41 B 32.837 49.537 -12.338 875 CF HIS 41 B 32.837 43.952 -12.505 876 O HIS 41 B 34.737 43.143 -11.564 877 N GLU 42 B 35.819 43.974 -13.373 878 CA GLU 42 B 36.805 42.897 -13.460 879 CB GLU 42 B 36.805 42.897 -13.460 887 CB GLU 42 B 36.806 41.769 -17.171 882 OE1 GLU 42 B 36.606 41.769 -17.171 882 OE1 GLU 42 B 36.606 41.776 -17.171 882 OE1 GLU 42 B 36.606 41.776 -17.171 882 OE1 GLU 42 B 36.806 41.776 -17.171 882 OE1 GLU 42 B 38.361 44.183 -12.561 887 CA LEU 43 B 39.538 41.815 -11.012 888 CB LEU 43 B 39.538 41.815 -11.012 889 CB LEU 43 B 39.970 42.470 -8.455 99.000 CD1 LEU 43 B 39.9673 32.100 -7.043 899 CG LEU 43 B 39.947 40.315 -11.580 899 CG LEU 43 B 39.947 40.315 -11.580 899 CG LEU 43 B 39.947 40.315 -11.580 899 CD1 TYR 44 B 41.804 36.795 -12.620 899 CD1 TYR 44 B 44.404 36.795 -12.620	THE LYS 40 B	858	N	LYS	40	В				
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899 CE1 11R 44 2	CEI 11R 44 D									
900 CD2 TYR 44 B 42.295 36.274 -13.634	CDZ TXR 44 B 42.233 35.274 13.031									

80/98 **Fig. 16\21** 

At	om	Resi	due	Chain	x	Y	z	δ
901	CE2	TYR	44	В	42.615	34.946	-13.694	0.56
902	CZ	TYR	44	В	43.862	34.534	-13.235	0.69
903	OH	TYR	44	В	44.205	33.222	-13.434	0.81
904	С	TYR	44	В	42.696	38.638	-10.395	0.51
905	0	TYR	44	В	43.481	39.582	-10.316	0.63
906	N	VAL	45	В	42.581	37.674	-9.534	0.53
907	CA	VAL	45	В	43.473	37.480	-8.368	0.53
908	CB	VAL	45	В	42.579	37.181	-7.148	0.56
909	CG1	VAL	45	В	43.320	36.968	-5.844	0.47
910	CG2	VAL	45	В	41.431	38.176	-7.035	0.52
911	C	VAL	45	В	44.398	36.280	-8.673	0.55
912	0	VAL	45	В	43.906	35.262	-9.193	0.53
913	N	SER	46	В	45.669	36.588	-8.765	0.59
914	CA	SER	46	В	46.829	35.718	-8.858	0.52
915	CB	SER	46	В	48.122	36.464	-9.287	0.46
916	OG	SER	46	В	48.841	35.585	-10.135	0.69
917	С	SER	46	В	47.138	35.163	-7.482	0.52
918	0	SER	46	В	47.307	35.955	-6.553	0.56
919	N	PHE	47	В	47.216	33.857	-7.380	0.59
920	CA	PHE	47	B	47.583	33.256	-6.069	0.57
921	CB	PHE	47	В	47.312	31.754	-6.088	0.53
922	CG	PHE	47	В	45.896	31.295	-6.174	0.55
923	CD1	PHE	47	В	44.869	32.001	-5.517	0.45
924	CD2	PHE	47	В	45.562	30.101	-6.818	0.54
925	CE1	PHE	47	В	43.554	31.589	-5.633	0.71
926	CE2	PHE	47	В	44.232	29.715	-7.010	0.56
927	CZ	PHE	47	В	43.221	30.426	-6.313	0.60
928	С	PHE	47	В	48.970	33.743	-5.648	0.49
929	0	PHE	47	В	49.222	33.901	-4.426	0.61
930	N	ARG	48	В	49.912	33.923	-6.541	0.51
931	CA	ARG	48	В	51.204	34.593	-6.314	0.57
932	CB	ARG	48	В	52.003	34.854	-7.564	0.49
933	CG	ARG	48	В	52.089	33.858	-8.662	0.74
934	CD	ARG	48	В	53.439	33.274	-8.844	0.83
935	NE	ARG	48	В	54.192	33.768	-9.964	0.71
936	CZ	ARG	48	В	54.800		-10.923	0.66
937	NH1	ARG	48	В	55.174		-10.831	0.68
938	NB2	ARG	48	· B	54.842	33.671	-12.123	0.61
939	С	ARG	48	<b>B</b>	51.122	35.885	-5.517	0.64
940	0	ARG	48	В	51.681	36.007	-4.397	0.69
941	N	ASP	49	В	50.284	36.811	-5.977	0.60
942	CA	ASP	49	В	49.970	38.044	-5.274	0.57
943	CB	ASP	49	В	48.961	38.897	-6.047	0.68
944	CG	ASP	49	В	49.395	39.128	-7.481	0.80
945	OD1	ASP	49	В	48.609	39.683	-8.272	0.89

81/98 *Fig. 16\22* 

A	tom	Resi	đuc	Chain	<u> </u>	<u>Y</u>	<u>z</u>	<u> </u>
		167	49	В	50.527	38.681	-7.801	0.83
946	OD2	ASP		В	49.619	37.883	-3.814	0.66
947	C	ASP	49	B	49.712	38.876	-3.062	0.72
948	0	ASP	49		48.879	36.830	-3.465	0.70
949	N	LEU	50	B B	48.428	36.605	-2.086	0.60
950	CA	LEU	50	B	47.115	35.800	-2.187	0.62
951	CB	LEU	50	В	46.024	36.479	-3.022	0.69
952	CG	LEU	50		44.677	35.940	-2.544	0.66
953	CD1	LEU	50	В		37.974	-2.667	0.66
954	CD2	LEU	50	В	46.100 49.471	35.808	-1.275	0.62
955	C	LEU	50	B B	49.097	35.367	-0.167	0.69
956	0	LEU	50 51	B	50.399	35.235	-2.005	0.62
957	N	GLY		В	51.320	34.191	-1.563	0.59
958	CA	GLY	51		50.554	32.900	-1.303	0.68
959	c	GLY	51	B B	50.408	32.521	-0.130	0.73
960	0	GLY	51	B	49.891	32.356	-2.325	0.60
961	N	TRP	52 52	B	49.127	31.117	-2.063	0.63
962	CA	TRP	52		47.650	31.299	-2.022	0.76
963	CB	TRP	52	В	46.961	32.057	-0.951	0.80
964	CG	TRP	52	B	45.593	32.517	-1.004	0.74
965	CD2	TRP	52	В	45.285	33.063	0.263	0.92
966	CE2	TRP	52	В	44.583	32.450	-1.963	1.01
967	CE3	TRP	52	В	47.376	32.267	0.333	0.82
968	CD1	TRP	52	В	46.402	32.927	1.050	0.73
969	NE1	TRP	52	B B	44.009	33.536	0.576	0.75
970	CZ2	TRP	52 52	<b>B</b>	43.332	32.969	-1.684	0.68
971	CZ3	TRP		В	43.053	33.511	-0.428	0.80
972	CH2	TRP	52	B	49.615	30.052	-3.046	0.72
973	C	TRP	52		49.190	28.891	-2.965	0.78
974	0	TRP	52	В	50.656	30.412	-3.769	0.64
975	N	GLN	53	B B	51.203	29.563	-4.822	0.74
976	CA	GLN	53		52.095	30.349	-5.798	0.68
977	CB	GLN	53	В	53.334	30.917	-5.138	0.5
978	CG	GLN	53	В	53.334	32.124	-4.268	0.7
979	CD	GLN	53	B	52.394	32.252	-3.363	0.7
980	OE1	GLN	53	В	54.224	33.011	-4.447	0.8
981	NE2	GLN	53	В	51.891	28.321	-4.280	0.7
982	C	GLN	53	В		27.451	-5.080	0.8
983	0	GLN	53	В	52.298	28.353	-3.040	0.7
984	N	ASP	54	B	52.335	27.288	-2.432	0.76
985	CA	ASP	54	В	53.153	27.716	-2.432	0.7
986	CB	ASP	54	В	53.499 54.614	28.739	-0.920	0.9
987	CG	ASP	54	В		28.739	-1.863	0.9
988	OD1	ASP	54	В	55.409			0.3
989	OD2	ASP	54	.В	54.784	29.235	0.217	
990	С	ASP	54	В	52.486	25.906	-2.491	0.8

Fig. 16\23

A	tom	Resi	duc	Chain	x	Y	z	<u> </u>
991	0	ASP	54	В	53.092	24.893	-2.898	0.82
992	N	TRP	55	B	51.285	25.839	-1.953	0.72
993	CA	TRP	55	В	50.495	24.644	-1.723	0.78
994	CB	TRP	55	В	49.926	24.753	-0.293	0.69
995	CG	TRP	55	В	51.035	24.962	0.691	0.80
996	CD2	TRP	55	В	52.396	24.529	0.465	0.76
997	CE2	TRP	55	В	53.149	24.987	1.569	0.85
998	CE3	TRP	55	В	52.924	23.554	-0.374	0.87
999	CD1	TRP	55	В	51.076	25.759	1.796	0.74
1000	NE1	TRP	55	В	52.345	25.793	2.337	0.76
1001	CZ2	TRP	55	В	54.479	24.626	1.745	0.78
1002	CZ3	TRP	55	В	54.216	23.134	-0.151	0.88
1003	CB2	TRP	55	В	54.968	23.639	0.905	0.82
1004	С	TRP	55	В	49.431	24.336	-2.770	0.84
1005	0	TRP	55	. В	48.906	23.197	-2.758	0.77
1006	N	ILE	56	В	49.105	25.287	-3.642	0.83
1007	CA	ILE	56	В	48.109	25.152	-4.693	0.78
1008	CB	ILE	56	В	47.382	26.490	-5.076	0.77
1009	CG2	ILE	56	<b>B</b>	47.094	26.569	-6.606	0.71
1010	CG1	ILE	56	В	46.030	26.567	-4.316	0.82
1011	CD	ILE	56	В	45.964	27.743	-3.298	0.90
1012	С	ILE	56	B	48.700	24.512	-5.944	0.74
1013	0	ILE	56	<b>B</b>	49.768	24.950	-6.388	0.80
1014	N	ILE	57	В	47.911	23.642	-6.567	0.68
1015	CA	ILE	57	В	48.281	22.985	-7.822	0.64
1016	CB	ILE	57	В	48.040	21.429	-7.725	0.65
1017	CG2	ILE	57	В	47.983	20.771	-9.109	0.55
1018	CG1	ILE	57	В	49.017	20.758	-6.736	0.64
1019	CD	ILE	57	В	48.780	19.230	-6.540	0.62
1020	С	ILE	57	В	47.565	23.593	-9.020	0.60
1021	0	ILE	57	В	48.093		-10.145	0.62
1022	N	ALA	58	В	46.279	23.911	-8.834	0.58
1023	CA	ALA	58	В	45.460	24.436	-9.967	0.49
1024	CB	ALA	58	В	44.970		-10.789	0.55
1025	C	ALA	58	В	44.202	25.091	-9.376	0.46
1026	0	ALA	58	В	43.929	24.534	-8.302	0.49
1027	N	PRO	59	В	43.872	26.310	-9.744	0.53
1028	CD	PRO	59	<b>B</b>	42.742	27.052	-9.192	0.46
1029	CA	PRO	59	B	44.401		-10.870	0.45
1030	CB	PRO	59	B	43.209		-11.429	0.47
1031	CG	PRO	59	B	42.321		-10.217	0.43
1032	С	PRO	59	B	45.527		-10.384	0.53
1033	0	PRO	59	В	46.032	27.913	-9.239	0.55
1034	N	GLU	60	В	46.018		-11.255	0.49
1035	CA	GLU	60	B	47.061	29.844	-10.776	0.58

# Fig. 16\24

		Resid	luc	Chain _	x	Υ	<b>Z</b>	δ
At	om	Acsic	iu C					
						<b>-</b> -		0.64
1036	СВ	GLU	60	В	48.099		-11.906	0.64
1037	CG	GLU	60	· <b>B</b>	49.287		-11.886	0.77
1038	CD	GLU	60	В	49.851		-13.082	0.79
1039	OE1	GLU	60	В	49.900		-13.170	0.84
1040	OE2	GLU	60	В	50.265		-13.997	0.61
1041	С	GLU	60	В	46.440		-10.374	0.53
1042	0	GLU	60	В	47.006	31.966	-9.605	0.52
1043	N	GLY	61	В	45.121		-10.496	0.52
1044	CA	GLY	61	В	44.275	32.336	-9.965	0.54
1045	С	GLY	61	В	42.888		-10.585	0.58
1046	0	GLY	61	В	42.568		-11.298	0.55
1047	N	TYR	62	В	42.036		-10.231	0.55
1048	CA	TYR	62	В	40.662		-10.845	0.54
1049	CB	TYR	62	В	39.780	32.410	-9.853	0.38
1050	CG	TYR	62	В	39.540	33.198	-8.558	0.47
1051	CD1	TYR	62	В	40.580	33.288	-7.649	0.47
1052	CE1	TYR	62	В	40.540	34.084		0.51
1053	CD2	TYR	62	В	38.335	33.785		0.45
1054	CE2	TYR	62	В	38.165	34.346		0.35
1055	cz	TYR	62	В	39.284	34.567		0.51
1056	OH	TYR	62	В	39.146	35.055		0.65
1057	С	TYR	62	В	40.133		-10.973	0.59
1058	0	TYR	62	В	40.590		-10.330	0.55
1059	N	ALA	63	В	39.076		-11.731	0.54
1060	CA	ALA	63	В	38.403		-11.984	0.49
1061	CB	ALA	63	В	37.619		-13,282	0.58
1062	С	ALA	63	В	37.422		-10.845	0.51
1063	0	ALA	63	В	36.491		-10.665	0.58
1064	N	ALA	64	В	37.780	37.157		0.54
1065	CA	ALA	64	B.	36.931	37.347		0.53
1066	СВ	ALA	64	В	37.812	37.741		0.53
1067	С	ALA	64	В	35.776	38.344		0.52
1068	0	ALA	64	В	34.882	38.380		0.54
1069	N	TYR	65	В	36.063	39.402		0.51
1070	CA	TYR	65	В	35.241	40.568		0.54
1071	CB	TYR	65	В	33.801		5 -10.438	0.47
1072	CG	TYR	65	В	33.841		1 -11.777	0.47
1073		TYR	65	В	34.042		5 ~12.974	0.44
1074		TYR	65	В	34.043		4 -14.190	0.47
1075		TYR	65	В	34.100		7 -11.756	0.61
1076		TYR	65	В	34.363		9 -12.922	0.44
1077		TYR		В	34.279		2 -14.139	0.67
1078		TYR		В	34.476		4 -15.253	0.70
1079		TYR		В	35.164	41.47		0.51
1080		TYR		В	35.488	40.94	3 -7.664	0.49

Fig. 16\25

Atom		Resi	duc	Chain	<u> </u>	Y	<u>z</u>	δ.
1081	ห	TYR	<b>6</b> 6	В	34.349	42.507	-8.928	0.52
1082	CA	TYR	66	В .	33.974	43.311	-7.729	0.50
1083	CB	TYP.	66	В	35.170	44.051	-7.126	0.60
1084	CG	TYR	66	В	35.700	45.179	-7.983	0.59
1085	CD1	TYR	66	в ,	36.704	44.935	-8.917	0.59
1086	CE1	TYR	66	В	37.279	45.951	-9.659	0.53
1087	CD2	TYR	66	В	35.376	46.512	-7.721	0.53
1088	CE2	TYR	66	В	36.057	47.548	-8.335	0.47
1089	CZ	TYR	66	В	36.836	47.261	-9.448	0.58
1090	OH	TYR	66	В	37.469		-10.066	0.70
1091	c	TYR	66	В	32.914	44.336	-8.166	0.51
1092	Õ	TYR	66	В	32.777	44.585	-9.351	0.52
1093	N	CYS	67	В	32.304	44.936	-7.173	0.58
1094	CA	CYS	67	В	31.148	45.843	-7.372	0.57
1095	C	CYS	67	В	31.597	47.265	-7.066	0.52
1096	Õ	CYS	67	В	32.160	47.475	-5.968	0.48
1097	CB	CYS	67	В	30.051	45.374	-6.413	0.54
1098	SG	CYS	67	В	29.272	43.787	-6.831	0.54
1099	N	GLU	68	В	31.226	48.194	-7.930	0.56
1100	CA	GLU	68	В	31.450	49.624	-7.617	0.47
1101	CB	GLU	68	В	32.888	50.026	-7.993	0.68
1102	CG	GLU	68	В	33.521	51.195	-7.264	0.53
1103	CD	GLU	68	В	34.985	51.442	-7.510	0.67
1104	OE1	GLU	68	В	35.681	52.049	-6.720	0.64
1105	OE2	GLU	68	B	35.357	51.239	-8.687	0.63
1106	C	GLU	6B	B	30.561	50.515	-8.485	0.43
1107	o	GLU	68	В	30.393	50.238	-9.663	0.42
1108	N	GLY	69	В	30.221	51.681	-7.922	0.48
1109	CA	GLY	69	В	29.452	52.707	-8.669	0.44
1110	C	GLY	69	- В .	28.406	53.234	-7.670	0.5
1111	o	GLY	69	В	28.223	52.696	-6.563	0.53
1112	N	GLU	70	В	27.714	54.297	-8.048	0.57
1113	CA	GLU	70	В	26.797	54.923	-7.116	0.6
1114	СВ	GLU	70	· <b>B</b>	26.834	56.401	-6.924	0.7
1115	CG	GLU	70	В	26.449	57.323	-8.062	0.7
1116	CD	GLU	70	В	26.041	58.691	-7.554	1.0
1117	OE1	GLU	70	В	26.122	58.985	-6.372	0.8
1118	OE2	GLU	70	В	25.470	59.352	-8.441	0.9
1119	C	GLU	70	В	25.392	54.363	-7.133	0.4
1120	Ö	GLU	70	В	24.945	53.921	-8.186	0.42
1121	N	CYS	71	В	24.785	54.485	-5.962	0.5
1122	CA	CYS	71	В	23.378	54.069	-5.805	0.6
1123	C	CYS	71	B	22.488	55.305	-5.637	0.5
1124	o	CYS	71	В	22.279	55.787	-4.547	0.5
1125	CB	CYS	71	В	23.223	52.994	-4.771	0.6
-143	CB	C13	, _	-		J	••••	•

85/98 **Fig. 16\26** 

At	om	Resi	iue	Chain	<u> </u>	<u>Y</u>	z	<u> </u>
								0 57
1126	SG	CYS	71	В	23.951	51.385	-5.196	0.57
1127	N	ALA	72	В	22.057	55.857	-6.745	0.50
1128	CA	ALA	72	В	21.341	57.102	-6.862	0.61
1129	CB	ALA	72	В	22.306	58.192	-7.425	0.61
1130	С	ALA	72	В	20.223	56.910	-7.903	0.62
1131	0	ALA	72	В	20.396	56.167	-8.880	0.59
1132	N	PHE	73	В	19.294	57.873	-7.785	0.59
1133	CA	PHE	73	В	18.244	57.978	-8.805	0.46
1134	CB	PHE	73	В	17.063	58.839	-8.472	0.54
1135	CG	PHE	73	В	16.335	58.428	-7.198	0.45
1136	CD1	PHE	73	В	15.725	57.183	-7.129	0.43
1137	CD2	PHE	73	В	16.303	59.266	-6.117	0.45
1138	CE1	PHE	73	B	15.178	56.702	-5.935	0.34
1139	CE2	PHE	73	В	15.627	58.907	-4.959	0.37
1140	cz	PHE	73	В	15.089	57.602	-4.895	0.33
1141	С	PHE	73	В	18.983	58.181	-10.083	0.52
1142	0	PHE	73	В	19.751	59.134	-9.925	0.54
1143	N	PRO	74	В.	18.492	-	-11.197	0.57
1144	ÇD	PRO	74	В	19.072		-12.495	0.57
1145	CA	PRO	74	В	17.638		-11.370	0.56
1146	CB	PRO	74	В	17.298		-12.832	0.52
1147	CG	PRO	74	В	18.376		-13.519	0.54
1148	С	PRO	74	В	18.219		-10.805	0.66
1149	0	PRO	74	В	19.370		-11.112	0.74
1150	N	LEU	75	В	17.359	54.601	-10.090	0.58
1151	CA	LEU	75	В	17.626	53.249	-9.627	0.61
1152	CB	LEU	75	В	17.151	53.088	-8.209	0.63
1153	CG	LEU	75	В	18.000	53.160	-7.006	0.60
1154	CD1	LEU	75	В	19.409	53.632	-7.025	0.57
1155	CD2	LEU	75	В	17.331	53.418	-5.703	0.49
1156	С	LEU	75	В	17.058		-10.726	0.68
1157	0	LEU	75	В	15.909		-10.655	0.72
1158	N	ASN	76	В	17.826		-11.802	0.68
1159	CA	ASN	76	В	17.569		-12.939	0.66
1160	CB	ASN	76	В	18.627		-14.028	0.62
1161	CG	ASN	76	В	18.661		-14.697	0.83
1162	OD1	ASN	76	В	17.657		-14.677	0.97
1163		ASN	76	В	19.757		-15.370	0.81
1164		ASN	76	В	17.288		-12.449	0.67
1165		ASN	76	В	17.535		-11.281	0.73
1166		SER	77	В	16.531		-13.260	0.73
1167		SER	77	В	16.114		-12.886	0.74
1168		SER	77	В	15.107		-13.894	0.54
1169		SER	77	В	14.248		-13.128	0.96
1170		SER	77	В	17.359	47.011	12.767	0.62

Fig. 16\27

A	tom	Resi	due	Chain	x	<u>Y</u>	Z	<u> </u>
				_				
1171	0	SER	77	В	17.477		-11.835	0.66
1172	N	TYR	78	В	18.200		-13.767	0.63
1173	CA	TYR	78	В	19.542		-13.850	0.75
1174	CB	TYR	78	В	20.249		-15.137	0.63
1175	CG	TYR	78	В	21.368	48.043	-14.976	0.91
1176	CD1	TYR	78	В	21.284	49.371	-15.447	0.81
1177	CE1	TYR	78	В	22.321	50.280	-15.228	0.88
1178	CD2	TYR	78	В	22.605	47.610	-14.494	0.98
1179	CE2	TYR	78	В	23.652	48.502	-14.265	0.99
1180	CZ	TYR	78	В	23.510	49.833	-14.640	0.94
1181	OH	TYR	78	В	24.611	50.630	-14.483	0.97
1182	С	TYR	78	В	20.376	46.79B	-12.592	0.7B
1183	0	TYR	78	В	21.428	46.139	-12.487	0.79
1184	N	MET	79	В	20.014	47.722	-11.719	0.74
1185	CA	MET	79	В	20.721	48.022	-10.475	0.62
1186	CB	MET	79	В	20.757	49.519	-10.191	0.68
1187	CG	MET	79	В	21.670	50.154	-11.207	0.51
1188	SD	MET	79	В	21.423	51.909	-11.149	0.67
1189	CE	MET	79	В	22.086	52.461	-9.625	0.53
1190	C	MET	79	В	20.193	47.231	-9.309	0.58
1191	0	MET	79	В	20.701	47.287	-8.186	0.61
1192	N	ASN	80	В	19.249	46.327	-9.575	0.55
1193	CA	ASN	80	В	18.791	45.510	-8.431	0.54
1194	CB	ASN	80	В	19.392	44.106	-8.469	0.86
1195	ÇG	ASN	80	В	19.052	43.259	-7.244	1.05
1196	OD1	ASN	80	В	19.403	43.560	-6.079	0.95
1197	ND2	ASN	80	В	18.498	42.064	-7.518	0.86
1198	С	ASN	80	В	18.879	46.202	-7.091	0.57
1199	0	ASN	80	В	19.195	45.517	-6.054	0.67
1200	N	ALA	81	В	17.955	47.146	-6.890	0.60
1201	CA	ALA	81	В	17.798	47.835	-5.577	0.52
1202	CB	ALA	81	В	17.298	49.261	-5.894	0.53
1203	С	ALA	81	В	16.837	46.990	-4.802	0.53
1204	0 -	ALA	81	В	15.878	46.552	-5.445	0.60
1205	N	THR	82	В	16.965	46.811	-3.500	0.54
1206	CA	THR	82	В	15.866	46.297	-2.680	0.44
1207	СВ	THR	82	В	16.477	45.801	-1.326	0.45
1208	0G1	THR	82	В	17.234	46.967	-0.821	0.49
1209	CG2	THR	82	В	17.599	44.729	-1.601	0.58
1210	C	THR	82	В	14.851	47.432	-2.443	0.67
1211	ō	THR	82	В	15.035	48.606	-2.852	0.55
1212	И	ASN	83	В	13.681	47.096	-1.913	0.61
1213	CA	ASN	83	В	12.741	48.177	-1.529	0.56
1214	CB	ASN	83	В	11.453	47.641	-0.901	0.53
1215	CG	ASN	В3	В	10.532	46.962	-1.900	0.48

Fig. 16\28

1216   OD1	A	tom	Resi	duc	Chain	<u>x</u>	Y	z	δ
1217 ND2					_	10 (50	47 100	2 100	0.55
1218   C									
1219 O ASN 83 B 13.074 50.239 -0.428 0.54 1220 N HIS 84 B 13.966 48.503 0.569 0.55 1221 CA HIS 84 B 14.811 49.226 1.518 0.41 1222 CB HIS 84 B 15.645 48.344 2.462 0.58 1223 CG HIS 84 B 16.315 49.085 3.572 0.40 1224 CD2 HIS 84 B 16.315 49.085 3.572 0.40 1224 CD2 HIS 84 B 17.7461 49.885 3.426 0.22 1226 CE1 HIS 84 B 17.727 50.393 4.612 0.18 1227 NE2 HIS 84 B 16.808 50.017 5.513 0.31 1229 O HIS 84 B 15.742 50.241 0.878 0.47 1229 O HIS 84 B 15.996 51.280 1.484 0.50 1230 N ALA 85 B 16.457 49.932 -0.190 0.52 1231 CA ALA 85 B 16.457 49.932 -0.190 0.52 1232 CB ALA 85 B 18.193 50.157 -1.868 0.48 1233 C ALA 85 B 16.658 52.059 -1.388 0.57 1234 O ALA 85 B 16.658 52.059 -1.388 0.57 1235 N ILE 86 B 15.570 51.726 -2.095 0.50 1236 CA ILE 86 B 17.384 53.017 -1.781 0.55 1237 CB ILE 86 B 13.487 52.116 -3.425 0.55 1238 CG2 ILE 86 B 13.487 52.116 -3.425 0.52 1239 CG1 ILE 86 B 13.972 51.293 -4.652 0.40 1240 CD ILE 86 B 12.867 53.315 -0.45 1242 O ILE 86 B 12.867 53.315 -0.618 0.49 1241 C ILE 86 B 12.867 53.315 -0.618 0.49 1242 C ILE 86 B 12.867 53.315 -0.618 0.49 1243 N VAL 87 B 13.880 53.315 -0.618 0.49 1244 CA VAL 87 B 13.880 53.315 -0.618 0.49 1245 CB VAL 87 B 13.507 54.142 0.517 0.45 1246 CG1 VAL 87 B 13.507 54.142 0.517 0.45 1247 CG2 VAL 87 B 13.507 54.422 0.517 0.43 1248 C VAL 87 B 13.507 54.422 0.517 0.43 1249 O VAL 87 B 13.507 54.422 0.517 0.43 1246 CG1 VAL 87 B 13.507 54.422 0.517 0.43 1247 CG2 VAL 87 B 13.507 54.422 0.517 0.43 1248 C VAL 87 B 13.507 54.422 0.517 0.43 1249 O VAL 87 B 13.507 54.422 0.517 0.43 1246 CG1 VAL 87 B 13.507 54.422 0.517 0.43 1247 CG2 VAL 87 B 13.507 54.422 0.517 0.45 1255 CB GLN 88 B 19.947 54.409 2.355 0.50 1256 NE2 GLN 88 B 19.947 54.409 2.355 0.50 1257 C GLN 88 B 19.947 54.409 2.355 0.50 1258 O GLN 88 B 19.947 54.409 2.355 0.50 1259 N TER 89 B 17.420 55.662 -0.420 0.52									
1220 N									
1221 CA									
1222 CB									
1223 CG									
1224 CD2 HIS 84 B 15.849 49.305 4.809 0.29 1225 ND1 HIS 84 B 17.461 49.885 3.426 0.22 1226 CE1 HIS 84 B 17.727 50.393 4.612 0.18 1227 NE2 HIS 84 B 17.727 50.393 4.612 0.18 1228 C HIS 84 B 15.742 50.241 0.878 0.47 1229 O HIS 84 B 15.742 50.241 0.878 0.47 1229 O HIS 84 B 15.996 51.280 1.484 0.50 1230 N ALA 85 B 16.457 49.932 -0.190 0.52 1231 CA ALA 85 B 16.457 49.932 -0.170 0.52 1233 C ALA 85 B 16.658 52.059 -1.388 0.57 1233 C ALA 85 B 18.193 50.157 -1.868 0.48 1233 C ALA 85 B 16.658 52.059 -1.388 0.57 1234 O ALA 85 B 17.384 53.017 -1.781 0.55 1235 N ILE 86 B 15.570 51.726 -2.095 0.50 1236 CA ILE 86 B 15.570 51.726 -2.095 0.50 1237 CB ILE 86 B 13.487 52.116 -3.425 0.57 1238 CG2 ILE 86 B 12.542 53.287 -3.912 0.42 1240 CD ILE 86 B 12.807 50.430 -52.234 0.49 1241 C ILE 86 B 12.807 50.430 -52.234 0.49 1242 C ILE 86 B 14.566 54.996 -1.894 0.58 1243 N VAL 87 B 13.880 53.315 -0.618 0.49 1244 CA VAL 87 B 13.880 53.315 -0.618 0.49 1244 CA VAL 87 B 13.880 53.315 -0.618 0.49 1245 CB VAL 87 B 13.507 54.142 0.517 0.42 1246 CG1 VAL 87 B 13.507 54.142 0.517 0.42 1247 CG2 VAL 87 B 13.507 54.142 0.517 0.42 1248 C VAL 87 B 12.892 53.242 1.570 0.43 1247 CG2 VAL 87 B 12.892 53.242 1.570 0.43 1249 O VAL 87 B 12.892 53.242 1.570 0.50 1250 N GLN 88 B 15.751 54.226 1.301 0.64 1251 CA GLN 88 B 15.751 54.226 1.301 0.64 1252 CB GLN 88 B 19.914 52.340 3.448 0.63 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1255 OE GLN 88 B 19.914 52.340 3.448 0.63 1255 OE GLN 88 B 19.914 52.340 3.448 0.63 1255 OE GLN 88 B 19.914 52.340 3.448 0.63									
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1240         CD         ILE         86         B         12.807         50.430         -5.234         0.49           1241         C         ILE         86         B         14.332         53.807         -1.751         0.45           1242         O         ILE         86         B         14.566         54.996         -1.894         0.58           1243         N         VAL         87         B         13.880         53.315         -0.618         0.49           1244         CA         VAL         87         B         13.507         54.142         0.517         0.42           1245         CB         VAL         87         B         12.781         53.866         2.950         0.39           1246         CG1         VAL         87         B         11.517         52.782         1.097         0.50           1248         C         VAL         87         B         11.517         52.782         1.097         0.50           1249         O         VAL         87         B         14.686         54.986         0.984         0.52           1249         O         VAL         87         B									
1241 C         ILE 86         B         14.332         53.807         -1.751         0.45           1242 O         ILE 86         B         14.566         54.996         -1.894         0.58           1243 N         VAL 87         B         13.880         53.315         -0.618         0.49           1244 CA         VAL 87         B         13.507         54.142         0.517         0.42           1245 CB         VAL 87         B         12.892         53.242         1.570         0.43           1246 CG1         VAL 87         B         12.781         53.866         2.950         0.39           1247 CG2         VAL 87         B         11.517         52.782         1.097         0.50           1248 C         VAL 87         B         14.686         54.986         0.984         0.52           1249 O         VAL 87         B         14.415         55.916         1.765         0.57           1250 N         GLN 88         B         15.751         54.226         1.301         0.64           1251 CA         GLN 88         B         17.007         54.802         1.773         0.50           1252 CB         GLN 88				-					
1242 O ILE 86 B 14.566 54.996 -1.894 0.58 1243 N VAL 87 B 13.880 53.315 -0.618 0.49 1244 CA VAL 87 B 13.507 54.142 0.517 0.42 1245 CB VAL 87 B 12.892 53.242 1.570 0.43 1246 CG1 VAL 87 B 12.781 53.866 2.950 0.39 1247 CG2 VAL 87 B 11.517 52.782 1.097 0.50 1248 C VAL 87 B 14.686 54.986 0.984 0.52 1249 O VAL 87 B 14.415 55.916 1.765 0.57 1250 N GLN 88 B 15.751 54.226 1.301 0.64 1251 CA GLN 88 B 17.007 54.802 1.773 0.50 1252 CB GLN 88 B 17.007 54.802 1.773 0.50 1253 CG GLN 88 B 19.477 54.409 2.355 0.50 1254 CD GLN 88 B 19.477 54.409 2.355 0.50 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1256 NE2 GLN 88 B 19.914 52.340 3.448 0.63 1258 O GLN 88 B 17.473 55.905 0.867 0.42 1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52									
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1245 CB VAL 87 B 12.892 53.242 1.570 0.43 1246 CG1 VAL 87 B 12.781 53.866 2.950 0.39 1247 CG2 VAL 87 B 11.517 52.782 1.097 0.50 1248 C VAL 87 B 14.686 54.986 0.984 0.52 1249 O VAL 87 B 14.415 55.916 1.765 0.57 1250 N GLN 88 B 15.751 54.226 1.301 0.64 1251 CA GLN 88 B 17.007 54.802 1.773 0.50 1252 CB GLN 88 B 18.035 53.927 2.372 0.54 1253 CG GLN 88 B 19.477 54.409 2.355 0.50 1254 CD GLN 88 B 20.342 53.445 3.126 0.53 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1256 NE2 GLN 88 B 19.914 52.340 3.448 0.63 1257 C GLN 88 B 17.473 55.905 0.867 0.42 1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52									
1246 CG1 VAL 87 B 12.781 53.866 2.950 0.39 1247 CG2 VAL 87 B 11.517 52.782 1.097 0.50 1248 C VAL 87 B 14.686 54.986 0.984 0.52 1249 O VAL 87 B 14.415 55.916 1.765 0.57 1250 N GLN 88 B 15.751 54.226 1.301 0.64 1251 CA GLN 88 B 17.007 54.802 1.773 0.50 1252 CB GLN 88 B 18.035 53.927 2.372 0.54 1253 CG GLN 88 B 19.477 54.409 2.355 0.50 1254 CD GLN 88 B 20.342 53.445 3.126 0.53 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1256 NE2 GLN 88 B 19.914 52.340 3.448 0.63 1257 C GLN 88 B 17.473 55.905 0.867 0.42 1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52									
1247         CG2         VAL         87         B         11.517         52.782         1.097         0.50           1248         C         VAL         87         B         14.686         54.986         0.984         0.52           1249         O         VAL         87         B         14.415         55.916         1.765         0.57           1250         N         GLN         88         B         15.751         54.226         1.301         0.64           1251         CA         GLN         88         B         17.007         54.802         1.773         0.50           1252         CB         GLN         88         B         18.035         53.927         2.372         0.54           1253         CG         GLN         88         B         19.477         54.409         2.355         0.50           1254         CD         GLN         88         B         20.342         53.445         3.126         0.53           1255         OE1         GLN         88         B         19.914         52.340         3.448         0.63           1256         NE2         GLN         88         B									
1248         C         VAL         87         B         14.686         54.986         0.984         0.52           1249         O         VAL         87         B         14.415         55.916         1.765         0.57           1250         N         GLN         88         B         15.751         54.226         1.301         0.64           1251         CA         GLN         88         B         17.007         54.802         1.773         0.50           1252         CB         GLN         88         B         18.035         53.927         2.372         0.54           1253         CG         GLN         88         B         19.477         54.409         2.355         0.50           1254         CD         GLN         88         B         20.342         53.445         3.126         0.53           1255         OE1         GLN         88         B         19.914         52.340         3.448         0.63           1256         NE2         GLN         88         B         21.607         53.766         3.183         0.42           1257         C         GLN         88         B         <									
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1251 CA GLN 88 B 17.007 54.802 1.773 0.50 1252 CB GLN 88 B 18.035 53.927 2.372 0.54 1253 CG GLN 88 B 19.477 54.409 2.355 0.50 1254 CD GLN 88 B 20.342 53.445 3.126 0.53 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1256 NE2 GLN 88 B 21.607 53.766 3.183 0.42 1257 C GLN 88 B 17.473 55.905 0.867 0.42 1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52		0	VAL	87					
1252 CB GLN 88 B 18.035 53.927 2.372 0.54 1253 CG GLN 88 B 19.477 54.409 2.355 0.50 1254 CD GLN 88 B 20.342 53.445 3.126 0.53 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1256 NE2 GLN 88 B 21.607 53.766 3.183 0.42 1257 C GLN 88 B 17.473 55.905 0.867 0.42 1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52	1250	N	GLN	88	В				
1253 CG GLN 88 B 19.477 54.409 2.355 0.50 1254 CD GLN 88 B 20.342 53.445 3.126 0.53 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1256 NE2 GLN 88 B 21.607 53.766 3.183 0.42 1257 C GLN 88 B 17.473 55.905 0.867 0.42 1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52	1251	CA	GLN	88	В				
1254 CD GLN 88 B 20.342 53.445 3.126 0.53 1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1256 NE2 GLN 88 B 21.607 53.766 3.183 0.42 1257 C GLN 88 B 17.473 55.905 0.867 0.42 1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52	1252	CB	GLN	88	В				
1255 OE1 GLN 88 B 19.914 52.340 3.448 0.63 1256 NE2 GLN 88 B 21.607 53.766 3.183 0.42 1257 C GLN 88 B 17.473 55.905 0.867 0.42 1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52	1253	CG	GLN	88					
1256 NE2 GLN 88 B 21.607 53.766 3.183 0.42 1257 C GLN 88 B 17.473 55.905 0.867 0.42 1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52	1254	CD	GLN	88					
1257 C GLN 88 B 17.473 55.905 0.867 0.42 1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52	1255	OE1	GLN	88					
1258 O GLN 88 B 18.004 56.920 1.314 0.52 1259 N THR 89 B 17.420 55.682 -0.420 0.52		NE2	GLN						
1259 N THR 89 B 17.420 55.682 -0.420 0.52	1257	С	GLN	88					
	1258	0	GLN						
1260 CA THR 89 B 17.748 56.663 -1.434 0.51	1259	N	THR						
	1260	CA	THR	89	В	17.748	56.663	-1.434	0.51

88/98 Fig. 16\29

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A	tom	Resi	đue	Chain	X	Y	Z	δ_
1261	СВ	THR	89	В	17.957	56.044	-2.830	0.44
1262	OG1	THR	89	В	18.892	54.925	-2.666	0.58
1263	CG2	THE	89	В	18.468	56, 959	-3.906	0.28
1264	С	THR	89	В	16.809	57.866	-1.437	0.60
1265	o	THR	89	В	17.292	58.957	-1.772	0.60
1266	N	LEU	90	В	15.600	57.756	-0.944	0.60
1267	CA	LEU	90	В	14.594	58.825	-0.965	0.54
1268	СВ	LEU	90	В	13.201	58.299	-1.267	0.64
1269	CG	LEU	90	В	12.008	59.230	-1.078	0.62
1270	CD1	LEU	90	В	11.967	60.201	-2.272	0.49
1271	CD2	LEU	90	В	10.728	58.385	-1.213	0.62
1272	c	LEU	90	В	14.771	59.667	0.279	0.47
1273	o	LEU	90	В	14.902	60.893	0.185	0.63
1274	N	VAL	91	В	15.098	59.037	1.376	0.43
1275	CA	VAL	91	В	15.498	59.640	2.618	0.40
1276	CB	VAL	91	В	15.662	58.617	3.726	0.35
1277	CG1	VAL	91	В	15.884	59.259	5.070	0.33
1278	CG2	VAL	91	В	14.368	57.821	3.941	0.47
1279	c	VAL	91	В	16.645	60.623	2.505	0.49
1280	ō	VAL	91	В	16.915	61.323	3.477	0.59
1281	N	HIS	92	В	17.546	60.291	1.635	0.51
1282	CA	HIS	92	В	18.848	60.835	1.409	0.56
1283	CB	HIS	92	В	19.806	59.782	0.770	0.55
1284	CG	HIS	92	В	21.133	60.406	0.469	0.52
1285	CD2	HIS	92	В	21.587	60.995	-0.668	0.36
1286	ND1	HIS	92	В	22.072	60.627	1.483	0.42
1287	CE1	HIS	92	В	23.116	61.193	0.876	0.49
1288	NE2	HIS	92	В	22.856	61.461	-0.379	0.43
1289	C	HIS	92	В	18.699	62.058	0.492	0.50
1290	ō	HIS	92	В	19.217	63.135	0.766	0.62
1291	N	PHE	93	В	17.866	61.925	-0.494	0.52
1292	CA	PHE	93	В	17.412	62.999	-1.335	0.62
1293	CB	PHE	93	В	16.605	62.527	-2.519	0.52
1294	CG	PHE	93	<b>B</b> .	15.862	63.671	-3.167	0.65
1295	CD1	PHE	93	B	14.542	63.901	-2.795	0.60
1296	CD2	PHE	93	В	16.473	64.466	-4.135	0.66
1297	CE1	PHE	93	В	13.873	65.043	-3.237	0.64
1298	CE2	PHE	93	- B	15.814	65.627	-4.580	0.66
1299	CZ	PHE	93	В	14.435	65.735	-4.295	0.51
1300	c	PHE	93	В	16.741	64.103	-0.506	0.66
1301	ō	PHE	93	В	17.155	65.264	-0.617	0.73
1302	N	ILE	94	В	15.788	63.768	0.315	0.64
1303	CA	ILE	94	В	15.077	64.619	1.252	0.61
1304	CB	ILE	94	В	13.867	63.896	1.897	0.63
1305	CG2	ILE	94	В	13.064	64.661	2.972	0.63

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Fig. 16\30

A	tom	Res	iduc	Chain	<u> </u>	Υ		δ
1306	CG1	ILE	94	В	12.891	63.334	0.814	0.51
1307	CD	ILE	94	В	11.846	62.434	1.602	0.63
1308	С	ILE	94	В	15.975	65.284	2.289	0.70
1309	0	ILE	94	В	15.535	66.310	2.824	0.75
1310	N	ASN	95	В	16.860	64.542	2.923	0.74
1311	CA	ASN	95	В	17.823	64.972	3.913	0.64
1312	CB	ASN	95	В	17.256	65.197	5.288	0.70
1313	CG	ASN	95	В	18.207	65.971	6.197	0.86
1314	OD1	ASN	95	В	19.322	66.347	5.782	0.79
1315	ND2	ASN	95	В	17.874	66.039	7.487	0.87
1316	C	ASN	95	В	19.132	64.178	3.867	0.65
1317	0	ASN	95	В	19.420	63.340	4.746	0.65
1318	N	PRO	96	В	20.049	64.736	3.096	0.64
1319	CD	PRO	96	В	19.759	65.911	2.228	0.68
1320	CA	PRO	96	В	21.354	64.191	2.805	0.65
1321	CB	PRO	96	В	22.034	65.260	1.943	0.69
1322	CG	PRO	96	В	20.869	65.837	1.180	0.72
1323	С	PRO	96	В	22.214	63.754	3.961	0.72
1324	0	PRO	96	В	23.152	62.938	3.812	0.73
1325	N	GLU	97	В	21.829	64.174	5.140	0.72
1326	CA	GLU	97	В	22.686	64.060	6.338	0.70
1327	CB	GLU	97	В	22.627	65.373	7.121	0.84
1328	CG	GLU	97	В	23.691	66.431	7.069	0.80
1329	CD	GLU	97	В	24.673	66.524	5.952	1.11
1330	OE1	GLU	97	В	25.881	66.300	6.044	1.13
1331	OE2	GLU	97	В	24.198	67.117	4.948	1.11
1332	С	GLU	97	В	22.108	62.957	7.224	0.72
1333	0	GLU	97	В	22.745	62.506	8.192	0.74
1334	N	THR	98	В	20.827	62.673	7.015	0.69
1335	CA	THR	98	В	20.159	61.643	7.830	0.67
1336	CB	THR	98	В	18.594	61.624	7.570	0.65
1337	OG1	THR	98	В	18.303	62.988	7.123	0.77
1338	CG2	THR	98	В	17.827	61.326	8.854	0.78
1339	С	THR	98	В	20.744	60.263	7.527	0.63
1340	0	THR	98	В	20.821	59.417	8.443	0.67
1341	N	VAL	99	В	21.028	60.046	6.249	0.57
1342	CA	VAL	99	В	21.514	58.705	5.846	0.52
1343	СВ	VAL	99	В	20.261	57.794	5.871	0.56
1344	CG1	VAL	99	В	19.489	57.979	4.573	0.44
1345	CG2	VAL	99	В	20.512	56.335	6.138	0.62
1346	С	VAL	99	В	22.242	58.813	4.532	0.52
1347	0	VAL	99	В	21.984	59.664	3.666	0.53
1348	N		100	В	23.279	57.962	4.403	0.53
1349	CD		100	В	23.638	56.903	5.352	0.54
1350	CA		100	В	23.966	57.811	3.125	0.49

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90/98 **Fig. 16\31** 

A	lom	Residuc	Chain	x	Y	<u>z</u>	
-							
1351	CB	PRO 100	В	25.101	56.822	3.411	0.50
1352	CG	PRO 100	В	24.727	56.112	4.666	0.47
1353	C	PRO 100	В	23.053	57.286	2.041	0.41
1354	0	PRO 100	В	22.099	56.537	2.250	0.58
1355	N	LYS 101	В	23.591	57.197	0.836	0.48
1356	CA	LYS 101	В	23.004	56.427	-0.241	0.40
1357	CB	LYS 101	В	23.525	56.940	-1.590	0.58
1358	CG	LYS 101	В	22.414	57.692	-2.367	0.69
1359	CD	LYS 101	В	22.889	58.782	-3.271	0.65
1360	CE	LYS 101	В	24.415	58.943	-3.297	0.61
1361	NZ	LYS 101	В	24.770	59.592	-4.589	0.76
1362	С	LYS 101	В	23.492	54.959	-0.048	0.56
1363	0	LYS 101	В	24.250	54.763	0.922	0.54
1364	N	PRO 102	В	22.672	54.006	-0.475	0.57
1365	CD	PRO 102	В	21.367	54.200	-1.164	0.48
1366	CA	PRO 102	В	23.080	52.604	-0.428	0.59
1367	СВ	PRO 102	В	21.934	51.832	-1.028	0.43
1368	CG	PRO 102	В	21.075	52.823	-1.736	0.54
1369	С	PRO 102	В	24.426	52.376	-1.076	0.5
1370	0	PRO 102	В	24.877	53.077	-2.006	0.63
1371	N	CYS 103	В	25.031	51.241	-0.744	0.58
1372	CA	CYS 103	В	26.294	50.811	-1.400	0.48
1373	CB	CYS 103	В	27.404	50.638	-0.384	0.75
1374	SG	CYS 103	В	27.517	48.995	0.344	0.83
1375	C	CYS 103	В	26.108	49.649	-2.355	0.5
1376	0	CYS 103	В	25.113	48.919	-2.353	0.53
1377	N	CYS 104	В	26.834	49.723	-3.461	0.32
1378	CA	CYS 104	В	26.958	48.724	-4.468	0.48
1379	С	CYS 104	В	27.702	47.477	-3.963	0.60
1380	0	CYS 104	В	28.863	47.528	-3.570	0.60
1381	СВ	CYS 104	В	27.694	49.397	-5.594	0.3
1382	SG	CYS 104	В	27.689	48.486	-7.124	0.5
1383	N	ALA 105	В	27.039	46.332	-4.024	0.5
1384	CA	ALA 105	В	27.391	45.157	-3.222	0.5
1385	СВ	ALA 105	В	26.840	45.289	-1.838	0.5
1386	c	ALA 105	В	26.862	43.953	-4.017	0.5
1387	o	ALA 105	В	26.240	44.236	-5.068	0.5
1388	N	PRO 106	В	27.562	42.850	-3.866	0.5
1389	CD	PRO 106	В	28.577	42.554	-2.822	0.5
1390	CA	PRO 106	В	27,352	41.676	-4.739	0.5
1391	CB	PRO 106	В	28.482	40.709	-4.321	0.5
1392	CG	PRO 106	В	29.464	41.531	-3.559	0.5
1393	C	PRO 106	В	25.996	41.079	-4.332	0.5
1394	0	PRO 106	В	25.637	41.105	-3.154	0.5
1305	N	TRO 100	R	25.222	40.677	-5.306	0.6

91/98 **Fig. 16\32** 

A	tom	Residue	Chain	X	<u>Y</u>		δ
		445	_		:		
1396	CA	THR 107	В	23.880	40.104	-4.913	0.65
1397	CB	THR 107	В	22.823	40.695	-5.928	0.45
1398	OG1	THR 107	В	23.160	40.092	-7.208	0.68
1399	CG2	THR 107	В	22.802	42.209	-6.091	0.56
1400	С	THR 107	В	24.011	38.593	-5.133	0.68
1401	0	THR 107	В	23.316	37.753	-4.543	0.68
1402	И	GLN 108	· B	25.040	38.240	-5.902	0.67
1403	CA	GLN 108	В	25.358	36.820	-6.067	0.64
1404	СВ	GLN 108	В	24.534	36.137	-7.105	0.58
1405	CG	GLN 108	В	24.889	36.327	-8.547	0.70
1406	CD	GLN 108	В	24.534	35.087	-9.358	0.99
1407	OE1	GLN 108	В	24.739	33.953	-8.916	0.83
1408	NE2	GLN 108	B	23.905	35.307	-10.508	0.94
1409	C	GLN 108	В	26.848	36.536	-6.135	0.63
1410	0	GLN 108	В	27.523	37.048	-7.047	0.64
1411	N	LEU 109	В	27.248	35.607	-5.288	0.60
1412	CA	LEU 109	В	28.574	34.966	-5.307	0.65
1413	CB	LEU 109	В	29.033	35.101	-3.831	0.55
1414	CG	LEU 109	В	29.122	36.560	-3.407	0.59
1415	CD1	LEU 109	В	28.855	36.704	-1.936	0.68
1416	CD2	LEU 109	В	30.501	37.120	-3.746	0.56
1417	С	LEU 109	В	28.563	33.511	-5.774	0.70
1418	0	LEU 109	B	27.843	32.695	-5.142 -6.507	0.75 0.67
1419	И	ASN 110	В	29.599	33.112	-6.849	
1420	CA	ASN 110	В	29.915	31.719	-8.350	0.65
1421	CB	ASN 110	В	30.080	31.445		0.59
1422	CG	ASN 110	B	28.797	31.687	-9.115	0.79
1423	OD1	ASN 110	В	28.778	31.854	-10.340	0.90
1424	ND2	ASN 110	В	27.734	31.706	-8.302	0.79
1425	C	ASN 110	В	31.162	31.203	-6.116	0.65
1426	0	ASN 110	В	32.020	31.945	-5.641	0.61
1427	N	ALA 111	В	31.238	29.877 29.116	-6.127 -5.489	0.68 0.64
1428	CA	ALA 111	В	32.307 31.831	27.798	-4.906	0.59
1429	CB	ALA 111	B	33.458	28.918	-6.467	0.57
1430	C	ALA 111	В	33.295	29.038	-7.697	0.54
1431	0	ALA 111	B B	34.629	28.709	-5.879	0.54
1432	N	ILE 112		35.738	28.160	-6.743	0.47
1433	CA	ILE 112	В		29.173	-6.985	0.47
1434	CB	ILE 112	В	36.886 36.603	30.359	-7.923	0.68
1435	CG2	ILE 112	В				0.56
1436	CG1	ILE 112	В	37.619	29.600	-5.712	
1437	CD	ILE 112	В	39.128 36.196	29.945 26.877	-5.888 -5.997	0.64
1438	C	ILE 112	<b>B</b>	36.186		-5.997	0.45 0.54
1439	0	ILE 112	В	36.185	26.857		
1440	N	SER 113	B	36.724	25.995	-6.770	0.44

92/98 Fig. 16\33

A	tom	Residuc	Chain	X	<u>Y</u>	<u>Z</u>	8
1441	CA	SER 113	В	37.490	24 802	_6 220	0.0
1442	CB	SER 113	В		24.802	-6.330	0.6
				36.951	23.581	-7.102	0.6
1443	OG C	SER 113	B	35.727	23.125	-6.515	0.6
1444		SER 113	В	38.962	25.003	-6.694	0.5
1445	0	SER 113	В	39.320	25.356	-7.828	0.52
1446	N	VAL 114	В	39.833	24.614	-5.778	0.5
1447	CA	VAL 114	В	41.265	24.576	-6.121	0.5
1448	CB	VAL 114	В	41.961	25.809	-5.514	0.6
1449	CG1	VAL 114	В	41.157	27.093	-5.667	0.49
1450	CG2	VAL 114	В	42.328	25.534	-4.094	0.5
1451	C	VAL 114	В	41.871	23.235	-5.686	0.60
1452	0	VAL 114	В	41.581	22.634	-4.637	0.5
1453	N	LEU 115	B	42.814	22.811	-6.510	0.49
1454	CA	LEU 115	В	43.490	21.511	-6.182	0.5
1455	CB	LEU 115	В	43.748	20.883	-7.603	0.55
1456	CG	LEU 115	B	44.210	19.423	-7.568	0.5
1457	CD1	LEU 115	B	43.121	18.518	-7.036	0.59
1458	CD2	LEU 115	B	44.655	19.035	-8.975	0.52
1459	C	LEU 115	B	44.739	21.880	-5.403	0.4
1460	0	LEU 115	В	45.582	22.571	-5.955	0.5
1461	N	TYR 116	В	44.954	21.369	-4.196	0.5
1462	CA	TYR 116	B	46.186	21.698	-3.479	0.69
L463	СВ	TYR 116	В	45.946	22.841	-2.480	0.70
1464	CG	TYR 116	B	45.134	22.457	-1.269	0.40
1465	CD1	TYR 116	B	43.762	22.395	-1.272	0.5
1466	CE1	TYR 116	В	43.062	21.927	-0.154	0.60
1467	CD2	TYR 116	В	45.764	22.284	-0.050	0.5
1468	CE2	TYR 116	В	45.120	21.791	1.067	0.52
469	CZ	TYR 116	В	43.748	21.620	1.016	0.70
1470	OH	TYR 116	В	43.110	21.288	2.184	0.78
471	С	TYR 116	В	46.810	20.454	-2.849	0.69
472	0	TYR 116	В	46.136	19.420	-2.752	0.69
	· N	PHE 117	В	48.067	20.582	-2.485	0.50
474	CA	PHE 117	В	48.891	19.633	-1.793	0.6
475	CB	PHE 117	В	50.185	19.204	-2.413	0.6
476	CG	PHE 117	В	51.022	20.135	-3.206	0.82
.477	CD1	PHE 117	В	51.097	20.008	-4.599	0.9
478	CD2	PHE 117	В	51.946	20.970	-2.561	1.13
479	CE1	PHE 117	В	51.960	20.804	-5.343	1.12
480	CE2	PHE 117	В	52.823	21.783	-3.292	1.03
481	CZ	PHE 117	В	52.837	21.686	-4.697	1.02
482	С	PHE 117	В	48.967	19.911	-0.315	0.62
483	0	PHE 117	В	49.562	20.918	0.059	0.72
484	N	ASP 118	В	48.287	19.095	0.474	0.67
485	CA	ASP 118	В	48.220	19.309	1.922	0.67

Fig. 16\34

A	tom	Residue	Chain	<u> </u>	Y	<u>z</u>	<u>δ</u>
		110	. 5	47 026	10 670	2.612	0 ~
1486	CB	ASP 118	'B	47.036	18.679	2.610	0.73
1487	CG	ASP 118	В	47.265	17.250	3.079	0.82
1488	OD1	ASP 118	В.	46.423	16.701	3.798	0.72
1489	OD2	ASP 118	В	48.369	16.720	2.831	0.60
1490	С	ASP 118	В	49.577	18.996	2.526	0.68
1491	0	ASP 118	В	50.536	18.660	1.826	0.75
1492	N	ASP 119	В	49.577	18.988	3.851	0.76
1493	CA	ASP 119	В	50.837	18.919	4.614	0.83
1494	CB	ASP 119	В	50.693	19.619	5.961	0.94
1495	CG	ASP 119	В	49.368	19.278	6.634	1.08
1496	OD1	ASP 119	В	49.132	18.110	6.991	1.04
1497	OD2	ASP 119	В	48.509	20.186	6.627	1.06
1498	С	ASP 119	В	51.341	17.482	4.673	0.87
1499	0	ASP 119	В	52.554	17.249	4.854	0.90
1500	N	SER 120	В	50.429	16.546	4.448	0.85
1501	CA	SER 120	В	50.821	15.123	4.389	0.80
1502	CB	SER 120	В	49.833	14.202	5.030	0.83
1503	OG	SER 120	В	48.833	14.876	5.781	1.02
1504	С	SER 120	В	51.188	14.778	2.957	0.80
1505	0	SER 120	В	51.840	13.749	2.686	0.83
1506	N	SER 121	В	50.860	15.712	2.064	0.73
1507	CA	SER 121	B	51.273	15.519	0.653	0.69
1508	CB	SER 121	В	52.529	14.649	0.607	0.66
1509	OG	SER 121	В	53.646	15.408	1.016	0.85
1510	C	SER 121	В	50.155	14.886	-0.163	0.63
1511	o	SER 121	В	50.340	14.556	-1.332	0.69
1512	N	ASN 122	В	49.022	14.741	0.477	0.65
1513	CA	ASN 122	В	47.764	14.427	-0.218	0.63
1514	CB	ASN 122	В	46.738	14.164	0.897	0.69
1515	CG	ASN 122	B	47.397	13.265	1.955	0.73
1516		ASN 122	В	47.766	12.140	1.597	0.73
1517	OD1 ND2	ASN 122	B	47.187	13.541	3.240	0.75
				47.412	15.553	-1.176	0.67
1518	C	ASN 122	В		16.728	-0.852	0.63
1519	0	ASN 122	В	47.620			
1520	N	VAL 123	В	46.918	15.189	-2.346	0.60
1521	CA	VAL 123	В	46.429	16.123	-3.360	0.6
1522	CB	VAL 123	B	46.774	15.543	-4.744	0.58
1523	CG1	VAL 123	B	45.814	16.141	-5.758	0.59
1524	CG2	VAL 123	В	48.224	15.828	-5.089	0.5
1525	С	VAL 123	В	44.917	16.290	-3.220	0.6
1526	0	VAL 123	В	44.237	15.263	-3.098	0.6
1527	N	ILE 124	В	44.491	17.451	-2.758	0.70
1528	CA	ILE 124	В	43.113	17.704	-2.302	0.65
1529	CB	ILE 124	В	43.231	18.240	-0.813	0.67
1530	CG2	ILE 124	В	41.850	18.575	-0.198	0.6

94/98 Fig. 16\35

A	tom	Re	sidue	Chain	<u>x</u>	Y	z	δ
1531	CG1	ILE	124	В	43.938	17.128	-0.011	0.66
1532	CD	ILE	124	В	43.524	16.995	1.468	0.84
1533	C	ILE	124	В	42.385	18.700	-3.208	0.67
1534	0	ILE	124	В	42.895	19.771	-3.5B2	0.63
1535	N	LEU	125	В	41.110	18.426	-3.448	0.63
1536	CA	LEU	125	В	40.235	19.393	-4.140	0.69
1537	CB	LEU	125	В	39.453	18.619	-5.187	0.67
1538	CG	LEU	125	В	38.491	19.404	-6.067	0.64
1539	CD1	LEU	125	В .	39.277	20.179	-7.130	0.49
1540	CD2	LEU	125	В	37.653	18.302	-6.750	0.48
1541	С	LEU	125	В	39.344	20.118	-3.135	0.67
1542	0	LEU	125	В	38.544	19.446	-2.470	0.66
1543	N	LYS	126	В	39.596	21.403	-2.961	0.66
1544	CA	LYS	126	В	38.865	22.238	-2.001	0.59
1545	CB	LYS	126	В	39.604	22.684	-0.788	0.61
1546	CG	LYS	126	В	38.871	23.673	0.128	0.78
1547	CD	LYS	126	В	39.041	23.334	1.601	0.91
1548	CE	LYS	126	В	38.196	24.169	2.543	0.82
1549	NZ	LYS	126	В	37.808	23.383	3.746	0.86
1550	С	LYS	126	В	38.054	23.325	-2.669	0.50
1551	0	LYS	126	В	38.253	23.700	-3.845	0.57
1552	N	LYS	127	В	36.810	23.373	-2.204	0.61
1553	CA		127	В	35.729	24.235	-2.664	0.60
1554	СВ		127	В	34.375	23.528	-2.620	0.58
1555	CG		127	В	33.312	24.232	-3.471	0.57
1556	CD		127	В	31.942	24.017	-2.833	0.62
1557	CE		127	В	30.852	24.324	-3.845	0.76
1558	NZ		127	В	31.445	24.255	-5.208	0.83
1559	С		127	В	35.626	25.471	-1.755	0.60
1560	0		127	В	35.815	25.364	-0.521	0.49
1561	N		128	В	35.798	26.599	-2.424	0.56
1562	CA		128	В	35.838	27.876	-1.669	0.50
1563	СВ		128	В	37.082	28.686	-2.089	0.77
1564	CG		128	В	38.294	28.241	-1.298	0.64
1565	CD1	TYR		B	39.395	27.674	-1.935	0.68
1566	CEI	TYR		В	40.468	27.198	-1.174	0.68
1567	CD2	TYR		В	38.204	28.154	0.085	0.72
1568	CE2	TYR		В	39.254	27.649	0.857	0.77
1569	CZ	TYR		B	40.332	27.060	0.200	0.89
1570	OH	TYR		В	41.314	26.484	0.961	0.90
1571	C	TYR		В	34.587	28.661	-2.110	0.46
1572	0	TYR		В	34.335	28.808	-3.308	0.59
1573	N	ARG		B	33.692	28.665	-3.308 -1.150	
1574	CA	ARG		В	32.380	29.311		0.56 0.61
1575	CB	ARG		В	31.479		-1.396	
		ARG	443		31.473	28.769	-0.273	0.74

95/98 Fig. 16\36

A	tom	Residuc	Chain	Х	Y	z	δ
1576	ÇG	ARG 129	В	31.060	27.303	-0.447	0.74
1577	CD	ARG 129	В	29.804	27.006	0.337	0.61
1578	NE	ARG 129	В	28.620	27.257	~0.488	1.06
1579	CZ	ARG 129	В	27.768	28,270	-0.310	1.07
1580	NH1	ARG 129	В	27.618	28.890	0.861	0.91
1581	NB2	ARG 129	В	27.093	28.771	-1.349	0.87
1582	С	ARG 129	В	32.587	30.829	-1.302	0.54
1583	0	ARG 129	В	33.426	31.314	-0.522	0.55
1584	N	ASN 130	В	31.860	31.561	-2.095	0.60
1585	CA	ASN 130	В	31.584	33.006	-2.013	0.59
1586	CB	ASN 130	В	31.088	33.375	-0.610	0.52
1587	CG	ASN 130	В	29.629	32.995	-0.392	0.63
1588	OD1	ASN 130	В	28.825	32.869	-1.318	0.58
1589	ND2	ASN 130	В	29.276	32.510	0.782	0.55
1590	С	ASN 130	В	32.864	33.763	-2.380	0.55
1591	0	ASN 130	В	33.205	34.674	-1.648	0.48
1592	N	MET 131	В	33.418	33.459	-3.520	0.54
1593	CA	MET 131	В	34.730	33.897	-3.974	0.51
1594	CB	MET 131	В	35.632	32.662	-4.327	0.54
1595	CG	MET 131	В	36.159	31.950	-3.137	0.44
1596	SD	MET 131	В	37.229	32.957	-2.052	0.61
1597	CE	MET 131	В	38.802	32.648	-2.915	0.60
1598	С	MET 131	В	34.538	34.709	-5.256	0.52
1599	0	MET 131	В	35.416	35.514	-5.570	0.59
1600	N	VAL 132	B	33.538	34.371	-6.055	0.59
1601	CA	VAL 132	В	33.347	35.038	-7.359	0.60
1602	CB	VAL 132	В	33.434	34.092	-8.517	0.52
1603	CG1	VAL 132	B	33.162	34.633	~9.898	0.48
1604	CG2	VAL 132	B	34.447	32.972	-8.485	0.66
1605	C	VAL 132	В	32.063	35.873	-7.308	0.69
1606	0	VAL 132	В	30.960	35.366	-7.012	0.63
1607	N	VAL 133	В	32.240	37.134	-7.668	0.65
1608	CA	VAL 133	В	31.100	38.075	-7.709	0.62
1609	СВ	VAL 133	В	31.530	39.550	-7.591	0.56
1610	CG1	VAL 133	В	30.408	40.468	-8.105	0.53
1611	CG2	VAL 133	В	31.852	39.841	-6.128	0.39
1612	C	VAL 133	B	30.409	37.812	-9.052	0.62
1613	0	VAL 133	B	31.105	38.041	-10.044	0.54
1614	N	ARG 134	B	29.134	37.398	-8.941	0.53
1615	CA	ARG 134	В	28.440	37.117	-10.208	0.56
1616	СВ	ARG 134	В	27.709	35.760	-10.158	0.68
1617	CG	ARG 134	В	28.433	34.636	-10.908	0.80
1618	CD	ARG 134	В	28.238	34.700	-12.379	0.79
1619	NE	ARG 134	В	27.810	33.460	-12.976	1.02
1620	cz	ARG 134	В	26.587	32.977	-13.159	1.10

Fig. 16\37

A	tom	Residue	Chain	x	Y	z	δ_
			_				
1621	NH1	ARG 134	В	25.494		-12.533	1.08
1622	NH2	ARG 134	В	26.450		-13.952	1.09
1623	C	ARG 134	В	27.517		-10.622	0.57
1624	0	ARG 134	В	27.185		-11.813	0.61
1625	N	ALA 135	В	27.038	39.030	-9.659	0.53
1626	CA	ALA 135	В	26.189	40.203	-9.986	0.49
1627	CB	ALA 135	В	24.713	39.740	-9.934	0.64
1628	С	ALA 135	В	26.352	41.156	-8.774	0.42
1629	0	ALA 135	В	26.515	40.639	-7.673	0.49
1630	N	CYS 136	B	25.926	42.365	-9.022	0.59
1631	CA	CYS 136	В	26.001	43.512	-8.089	0.67
1632	С	CYS 136	В	24.646	44.198	-7.969	0.61
1633	0	CYS 136	В	23.917	44.273	-8.985	0.59
1634	CB	CYS 136	В	26.990	44.534	-8.701	0.61
1635	SG	CYS 136	В	28.689	43.852	-8.673	0.61
1636	N	GLY 137	В	24.362	44.756	-6.808	0.59
1637	CA	GLY 137	В	23.099	45.570	-6.772	0.59
1638	С	GLY 137	В	23.290	46.706	-5.787	0.52
1639	0	<b>GLY 137</b>	В	24.077	46.520	-4.845	0.49
1640	N	CYS 138	В	22.261	47.515	-5.634	0.57
1641	CA	CYS 138	В	22.248	48.539	-4.545	0.48
1642	С	CYS 138	В	21.475	48.015	-3.369	0.56
1643	0	CYS 138	В	20.235	47.865	-3.446	0.55
1644	CB	CYS 138	В	21.556	49.798	-5.123	0.38
1645	SG	CYS 138	В	22.664	50.498	-6.428	0.57
1646	N	HIS 139	В	22.137	48.025	-2.210	0.63
1647	CA	HIS 139	B	21.436	47.660	-0.970	0.54
1648	CB	HIS 139	В	20.837	46.284	-0.842	0.88
1649	CG	HIS 139	В	21.539	45.272	-1.707	0.94
1650	CD2	HIS 139	В	22.879	45.046	-1.811	0.79
1651	ND1	HIS 139	В	20.936	44.549	-2.706	0.91
1652	CE1	HIS 139	В	21.855	43.796	-3.288	0.97
1653	NE2	HIS 139	В	23.034	44.089	-2.774	0.96
1654	C	HIS 139	В	21.888	48.359	0.264	0.58
1655	OT1	HIS 139	В	21.160	48.202	1.268	0.75
1656	OT2	HIS 139	В	22.865	49.139	0.256	0.79
1657	OT	WAT 201	A	31.351	45.516	-2.695	0.57
165B	OT	WAT 202	A	10.574	42.304	-3.269	0.77
1659	OT	WAT 203	A	41.094	47.385	-10.715	0.74
1660	OT	WAT 204	A	-6.527	46.271	-4.416	0.83
1661	OT	WAT 205	A	-7.389	42.963	-2.480	0.75
1662	OT	WAT 206	A	-5.998	42.514	2.104	0.60
1663	OT	WAT 207	A	25.154	37.549	3.436	0.76
1664	OT	WAT 208	A	31.925	33.286	2.732	0.58
1665	OT	WAT 209	A	32.701	43.734	-4.779	0.52

Fig. 16\38

A	lom	Residue	Chain	x	Y	z	δ_
1666	OT	WAT 210	A	15.485	51.948	13.181	0.60
1667	OT	WAT 211	A	9.829	38.538	6.111	0.71
1668	OT	WAT 212	A	11.550	40.302	3.050	0.58
1669	OT	WAT 213	A	42.134	46.885	5.506	0.73
1670	OT	WAT 214	A	37.738	52.318	-0.352	0.72
1671	OT	WAT 215	A	40.582	52.333	-4.324	0.66
1672	OT	WAT 216	A	22.375	54.496	15.975	0.63
1673	OT	WAT 217	A	49.983	39.399	2.310	0.69
1674	OT	WAT 218	A	5.369	58.756	12.045	0.61
1675	OT	WAT 219	A	0.867	40.439	5.311	0.67
1676	OT	WAT 220	A	25.522	37.902	-0.824	0.80
1677	OT	WAT 221	A	12.228	59.495	7.513	0.81
1678	OT	WAT 222	A	10.798	47.556	11.898	0.67
1679	OT	WAT 223	A	0.494	40.963	0.254	0.75
1680	OT	WAT 224	A	33.591	41.614	-1.644	0.50
1681	OT	WAT 225	A	24.730	59.387	8.279	0.74
1682	OT	WAT 226	A	17.020	38.835	3.348	0.73
1683	OT	WAT 227	A	34.395	49.780	3.674	0.61
1684	OT	WAT 228	A	6.972	43.390	-2.094	0.59
1685	OT	WAT 229	A	25.493	43.453	12.008	0.68
1686	OT	WAT 230	A	31.349	49.756	-3.421	0.70
1687	OT	WAT 231	A	2.519	49.133	-4.219	0.80
1688	OT	WAT 232	A	24.405	52.256	3.441	0.61
1689	OT	WAT 233	A	-0.363	65.457	3.148	0.79
1690	OT	WAT 201	В	23.742	49.909	2.695	0.57
1691	OT	WAT 202	В	31.349	30.309	3.269	0.77
1692	OT	WAT 203	В	20.489	59.281	10.715	0.74
1693	OT	WAT 204	В	43.335	17.483	4.416	0.83
1694	OT	WAT 205	В	40.901	15.082	2.480	0.75
1695	OT	WAT 206	В	39.817	16.063	-2.104	0.60
1696	OT	WAT 207	В	19.941	40.558	-3.436	0.76
1697	OT	WAT 208	В	12.864	44.291	-2.732	0.58
1698	OT	WAT 209	В	21.524	50.187	4:779	0.52
1699	OT	WAT 210	В	37.246	39.384	-13.181	0.60
1700	OT	WAT 211	В	28.460	27.781	-6.111	0.71
1701	OT	WAT 212	В	29.127	30.154	-3.050	0.58
1702	OT	WAT 213	В.	19.536	59.931	-5.506	0.73
1703	OT	WAT 214	В	26.439	58.841	0.352	0.72
1704	OT	WAT 215	В	25.030	61.311	4.324	0.66
1705	OT	WAT 216	В	36.007	46.625	-15.975	0.63
1706	OT	WAT 217	В	9.129	62.986	-2.310	0.69
1707	OT	WAT 218	В	48.199	34.028	-12.045	0.61
1708	OT	WAT 219	В	34.587	20.970	-5.311	0.67
1709	OT	WAT 220	В	20.063	41.054	0.824	0.80
1710	OT	WAT 221	В	45.410	40.337	-7.513	0.81
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98/98 **Fig. 16\39** 

Atom		Res	due Chain	<u> </u>	<u> Y</u>	<u>z</u>	<u> </u>	
1711	OT	WAT	222	В	35.785	33.129	-11.898	0.67
1712	OT	WAT	223	В	35.228	20.909	-0.254	0.75
1713	OT	WAT	224	В	19.243	49.897	1.644	0.50
1714	OT	TAW	225	В	39.065	51.110	-8.279	0.74
1715	OT	WAT	226	B	25.122	34.157	-3.348	0.73
1716	OT	WAT	227	В	25.913	54.677	-3.674	0.61
1717	OT	WAT	228	В	34.091	27.733	2.094	0.59
1718	OT	WAT	229	В	24.885	43.804	-12.008	0.68
1719	OT	WAT	230	В	27.415	52.027	3.421	0.70
1720	OT	WAT	231	В	41.291	26.748	4.219	0.80
1721	OT	WAT	232	В	33.052	47.263	-3.441	0.61
1722	OT	WAT	233	В	56.869	32.414	-3.148	0.79

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## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:		(11) International Publication Number: WO 97/26277
C07K 14/51, G06F 17/50, C07K 1/00	A3	(43) International Publication Date: 24 July 1997 (24.07.97)
(21) International Application Number: PCT/US9 (22) International Filing Date: 22 January 1997 (2)		CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
<ul> <li>(30) Priority Data: 08/589,552 22 January 1996 (22.01.96)</li> <li>(71) Applicants: CREATIVE BIOMOLECULES, INC. [ 45 South Street, Hopkinton, MA 01748 (US). BRA UNIVERSITY [US/US]; 45 South Street, Waltha 02254-9110 (US).</li> <li>(72) Inventors: KECK, Peter, 50 Dolan Road, Millbury, M. (US). GRIFFITH, Diana, L.; 15 Woodridge Circle, MA 02193 (US). CARLSON, William, D.; 40 Bl. Road, Weston, MA 02193 (US). RUEGER, David Downey Street, Hopkinton, MA 01748 (US). SAN Kuber, T.; 6 Spring Street, Medway, MA 02053 (U)</li> <li>(74) Agent: GREENHALGH, Duncan, A.; Testa, Hur Thibeault, L.L.P., High Street Tower, 125 High Boston, MA 02110 (US).</li> </ul>	ANDEI AM, M. A 0152 Westor ack Oa I, C.; 1 MPATI- JS).	Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.  (88) Date of publication of the international search report:  25 September 1997 (25.09.97)

## (54) Title: MORPHOGEN ANALOGS AND METHODS FOR PRODUCING THEM

#### (57) Abstract

The invention disclosed herein provides methods and compositions for the computer-assisted design of morphogen analogs. Practice of the invention is enabled by the use of at least a portion of the atomic co-ordinates defining the three-dimensional structure of human osteogenic protein-1 (hOP-1) as a starting point in the design of the morphogen analogs. In addition, the invention provides methods for producing morphogen analogs of interest, and methods for testing whether the resulting analogs mimic or agonize human OP-1-like biological activity. The invention also provides a family of morphogen analogs produced by such methods.

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Inten 1al Application No PCT/US 97/01071

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X	WO 89 09788 A (CREATIVE BIOMOLEC October 1989 see claims 21-42	22,23					
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